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(54) Title: METHOD AND REAGENT FOR INHIBITING CANCER DEVELOPMENT

(57) Abstract

An enzymatic RNA molecule which specifically cleaves mRNA encoded by an mdr-1 gene, or associated with development or maintenance of chronic myelogenous leukemia, promyelocytic leukemia, Burkitt's lymphoma or acute lymphocytic leukemia, follicular lymphoma, B-cell acute lymphocytic leukemia, breast cancer, colon carcinoma, neuroblastoma, and lung cancer.

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DESCRIPTION

METHOD AND REAGENT FOR INHIBITING CANCER DEVELOPMENT

Background of the Invention

This invention relates to methods for treating cancer, and in particular, growth of a transformed cell, and inhibition of progression to a transformed phenotype in pre-neoplastic cells.

Transformation is a cumulative process whereby normal control of cell growth and differentiation is interrupted, usually through the accumulation of mutations affecting the expression of genes that regulate cell growth and differentiation.

Scanlon W091/18625, W091/18624, and W091/18913 describes a ribozyme effective to cleave oncogene RNA in the H-ras gene. This ribozyme is said to inhibit C-fos expression in response to cis-plantin or other stimuli. Reddy, WO 92/00080 and U.S. Serial No. 07/544,199 (filed June 26, 1990), describes use of ribozymes as therapeutic agents for leukemias, such as chronic myelogenous leukemia (CML) by targeting specific junction regions of the bcr-abl fusion transcript.

20 Summary of the Invention

This invention concerns use of a ribozyme targeted to the P-glycoprotein (mdr-1 gene) or other cancer-related genes prior to and/or during administration of anticancer chemotherapeutic agents. Inclusion of such a ribozyme increases the susceptibility of the transformed cells to such agents.

Applicant notes that relapse of disease caused by cancerous cells after administration of chemotherapeutic agents is a major problem in obtaining lasting remissions in a clinic. In some neoplasias, relapse is caused by the expansion of a population of transformed cells resistant to the initial and subsequent

forms of chemotherapy due to inappropriate expression of the mdr-1 gene, also called P-glycoprotein. expression is usually caused by selection of transformed cells that have amplified the mdr-1 gene and thus produce 5 increased amounts of the mdr-1 gene product. describes treatment of and prevention of this condition by use of ribozymes targeted to the mRNA encoded by this gene.

The mdr-1 gene encodes a 170 kDa integral membrane transport protein that confers resistance to certain chemotherapeutic agents such as colchicine, doxorubicin, actinomycin D and vinblastine (reviewed in Gottesman and Pastan, 263 J. Biol. Chem. 12163, 1988). The gene has been isolated from both human and rodent 15 cells selected in vitro for resistance to such agents (Roninson et al., 309 Nature 626, 1984; and Roninson et al., 83 Proc. Natl. Acad. Sci USA, 4538, 1986), and the entire 4.5-kb MDR1 transcript encoding the human MDR1 has been sequenced (Chen et al., 47 Cell 381, 1986, EMBL 20 accession # M14758). The gene is normally expressed in the cells of the colon, small intestine, kidney, liver and adrenal gland. High levels of MDR1 transcript have been found in adenocarcinomas that are intrinsically resistant to a broad range of chemotherapeutic agents, such as those 25 derived from adrenal, kidney, liver and bowel.

The invention features use of ribozymes to inhibit the development or expression of a transformed phenotype in man and other animals by modulating expression of a gene that either contributes to, or inhibits the expression of CML, promyelocytic leukemia, Burkitt's lymphoma, acute lymphocytic leukemia, follicular lymphoma, B-cell acute lymphocytic leukemia, breast cancer, colon carcinoma, neuroblastoma, lung cancer, and other neoplastic conditions. Cleavage of targeted mRNAs 35 expressed in pre-neoplastic and transformed cells elicits inhibition of the transformed state.

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Ribozymes are RNA molecules having an enzymatic activity which is able to repeatedly cleave other separate RNA molecules in a nucleotide base sequence specific Such enzymatic RNA molecules can be targeted to 5 virtually any RNA transcript and efficient cleavage has been achieved in vitro. Kim et al., 84 Proc. Natl. Acad. Sci. USA 8788, 1987; Haseloff and Gerlach, 334 Nature 585, 1988; Cech, 260 <u>JAMA</u> 3030, 1988; and Jefferies et al., 17 Nucleic Acids Research 1371, 1989.

Ribozymes act by first binding to a target RNA. Such binding occurs through the target RNA binding portion of a ribozyme which is held in close proximity to an enzymatic portion of the RNA which acts to cleave the target RNA. Thus, the ribozyme first recognizes and then binds a target RNA through complementary base-pairing, and once bound to the correct site, acts enzymatically to cut the target RNA. Strategic cleavage of such a target RNA will destroy its ability to direct synthesis of an encoded protein. After a ribozyme has bound and cleaved its RNA 20 target it is released from that RNA to search for another target and can repeatedly bind and cleave new targets.

ribozyme The enzymatic nature of a advantageous over other technologies, such as antisense technology (where a nucleic acid molecule simply binds to 25 a nucleic acid target to block its translation) since the effective concentration of ribozyme necessary to effect a therapeutic treatment is lower than that of an antisense. This advantage reflects the ability of oligonucleotide. a single the ribozyme to act enzymatically. Thus, 30 ribozyme molecule is able to cleave many molecules of In addition, the ribozyme is a highly target RNA. specific inhibitor, with the specificity of inhibition depending not only on the base pairing mechanism of binding, but also on the mechanism by which the molecule inhibits the expression of the RNA to which it binds. That is, the inhibition is caused by cleavage of the RNA target and so specificity is defined as the ratio of the rate of cleavage of the targeted RNA over the rate of cleavage of non-targeted RNA. This cleavage mechanism is dependent upon factors additional to those involved in base pairing. Thus, it is thought that the specificity of action of a ribozyme is greater than that of antisense oligonucleotide binding the same RNA site.

This class of chemicals exhibits a high degree of specificity for cleavage of the intended target mRNA. Consequently, the ribozyme agent will only affect cells expressing that particular gene, and will not be toxic to normal tissues.

Thus, the invention features novel enzymatic RNA molecules, or ribozymes, and methods for their use for inhibiting cancer-related mRNA expression. Such ribozymes 15 can be used in a method for treatment of disease caused by expression of the cancer-related genes in man and other animals, including other primates. This conclusion, as noted above, is based upon the finding that many forms of cancer become unresponsive to certain chemotherapeutic agents as a result of overexpression of, e.g., the mdr-1 The advantage of using ribozymes of the present invention is their ability to specifically cleave the targeted mRNA, ultimately leading to a reduction in target gene activity through a decrease in level of the gene product. Use of mdr-1 specific ribozymes removes the mechanism of drug resistance used by transformed cells, and thus enhances drug therapies for tumor cell growth. These agents can be administered prior to and during chemotherapeutic treatment of those neoplasias known to have a high incidence of drug resistance, or can be used prophylactically for all neoplasias.

The invention can also be used to treat cancer or pre-neoplastic conditions. Two preferred administration protocols can be used, either in vivo administration to reduce the tumor burden, or ex vivo treatment to eradicate transformed cells from tissues such as bone marrow prior to reimplantation.

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Thus, in a first aspect, the invention features an enzymatic RNA molecule (or ribozyme) which cleaves mdr-1 mRNA (i.e., mRNA expressed from the mdr-1 gene), or its In particular, the invention features equivalent. 5 hammerhead ribozymes designed to cleave accessible areas Such areas include those sequences of the mdr-1 mRNA. shown in Fig. 2.

In a second aspect, the invention features an enzymatic RNA molecule (or ribozyme) which cleaves mRNA associated with development or maintenance of CML, promyelocytic leukemia, Burkitt's lymphoma or lymphocytic leukemia, follicular lymphoma, B-cell acute lymphocytic leukemia, breast cancer, colon carcinoma, neuroblastoma, and lung cancer, including mRNA targets disclosed in Figs. 3 to 11. Such mRNA is recognized by those in the art to encode an aberrant cellular protein which is able to control cellular proliferation, and is directly linked to (correlated with) the presence of the leukemic phenotype.

By "enzymatic RNA molecule" it is meant an RNA molecule which has complementarity in a substrate binding region to a specified mRNA target, and also has an enzymatic activity which is active to specifically cleave RNA in that mRNA. That is, the enzymatic RNA molecule is 25 able to intermolecularly cleave mRNA and thereby inactivate a target mRNA molecule. This complementarity functions to allow sufficient hybridization of enzymatic RNA molecule to the target RNA to allow the cleavage to occur. For in vivo use, such complementarity 30 may be between 30 and 45 bases. One hundred percent complementarity is preferred, but complementarity as low as 50-75% may also be useful in this invention.

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By "equivalent" RNA to mdr-1 mRNA is meant to mRNA molecules naturally occurring include those associated with neoplastic diseases in various animals, including humans, and other primates, which have similar structures and functions to that mdr-1 mRNA in humans.

The deduced sequences of the mouse and human Pglycoproteins are 80% identical.

In preferred embodiments, the enzymatic RNA molecule is formed in a hammerhead motif, but may also be formed in the motif of a hairpin, hepatitis delta virus, group I intron or RNaseP-like RNA (in association with an RNA guide sequence). Examples of such hammerhead motifs are described by Rossi et al., 8 Aids Research and Human Retroviruses 183, 1992; of hairpin motifs by Hampel et 10 al., "RNA Catalyst for Cleaving Specific RNA Sequences", filed September 20, 1989, which is a continuation-in-part of U.S. Serial No. 07/247,100 filed September 20, 1988, Hampel and Tritz, 28 Biochemistry 4929, 1989 and Hampel et al., 18 Nucleic Acids Research 299, 1990; an example of 15 the hepatitis delta virus motif is described by Perrotta and Been, 31 Biochemistry 16, 1992; of the RNaseP motif by Guerrier-Takada et al., 35 Cell 849, 1983; and of the group I intron by Cech et al., U.S. Patent 4,987,071. These specific motifs are not limiting in the invention 20 and those skilled in the art will recognize that all that is important in an enzymatic RNA molecule of this invention is that it has a specific substrate binding site which is complementary to one or more of the target mRNA regions, and that it have nucleotide sequences within or 25 surrounding that substrate binding site which impart an mRNA cleaving activity to the molecule.

In a related aspect, the invention features a mammalian cell which includes an enzymatic RNA molecule as described above. Preferably, the mammalian cell is a 30 human or other primate cell.

In another related aspect, the invention features an expression vector which includes nucleic acid encoding the enzymatic RNA molecules described above, located in the vector, <u>e.g.</u>, in a manner which allows expression of that enzymatic RNA molecule within a mammalian cell.

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In yet another related aspect, the invention features a method for treatment of an mdr-1 gene-related disease, chronic myelogenous leukemia (CML), promyelocytic leukemia, Burkitt's lymphoma or acute lymphocytic leukemia, follicular lymphoma, B-cell acute lymphocytic leukemia, breast cancer, colon carcinoma, neuroblastoma, or lung cancer, by administering to a patient an enzymatic RNA molecule as described above.

In another related aspect, the invention 10 features a method for treatment of CML by ex vivo treatment of blood or marrow cells with an enzymatic RNA molecule as described above.

The invention provides a class of chemical cleaving agents which exhibit a high degree of specificity 15 for the mRNA causative of CML, promyelocytic leukemia, Burkitt's lymphoma or acute lymphocytic leukemia, follicular lymphoma, B-cell acute lymphocytic leukemia, breast cancer, colon carcinoma, and neuroblastoma. desired, such ribozymes can be designed to target equivalent single-stranded DNAs by methods known in the art. The ribozyme molecule is preferably targeted to a highly conserved sequence region of the mdr-1 mRNA. Such enzymatic RNA molecules can be delivered exogenously to affected cells or endogenously to infected cells. 25 preferred hammerhead motif the small size (less than 40 nucleotides, preferably between 32 and 36 nucleotides in length) of the molecule allows the cost of treatment to be reduced compared to other ribozyme motifs.

The smallest ribozyme delivered for any type of treatment reported to date (by Rossi et al., 1992, supra) is an in vitro transcript having a length of 142 nucleotides. Synthesis of ribozymes greater than 100 nucleotides in length is very difficult using automated methods, and the therapeutic cost of such molecules is prohibitive. Delivery of ribozymes by expression vectors is primarily feasible using only ex vivo treatments. This limits the utility of this approach. In this invention,

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small ribozyme motifs (e.g., of the hammerhead structure, shown generally in Fig. 1) are used for exogenous delivery. The simple structure of these molecules also increases the ability of the ribozyme to invade targeted 5 regions of the mRNA structure. Thus, unlike the situation when the hammerhead structure is included within longer transcripts, there are no non-ribozyme flanking sequences to interfere with correct folding of the ribozyme structure, as well as complementary binding of the ribozyme to the mRNA target.

The enzymatic RNA molecules of this invention can be used to treat human CML, promyelocytic leukemia, Burkitt's lymphoma, acute lymphocytic leukemia, follicular lymphoma, B-cell acute lymphocytic leukemia, breast 15 cancer, or lung cancer. Affected animals can be treated at the time of cancer, or in a prophylactic manner. This timing of treatment will reduce the number of affected cells and disable cellular replication. This is possible because the ribozymes are designed to disable those structures required for successful cellular proliferation.

Ribozymes of this invention may be used as diagnostic tools to examine genetic drift and mutations The close relationship between within diseased cells. ribozyme activity and the structure of the target RNA 25 allows the detection of mutations in any region of the molecule which alters the base-pairing and threedimensional structure of the target RNA. By using multiple ribozymes described in this invention, one may map nucleotide changes which are important to 30 structure and function in vitro, as well as in cells and tissues. Cleavage of target RNAs with ribozymes may be used to inhibit gene expression and define the role specified gene products in (essentially) of progression of disease. In this manner, other genetic targets may be defined as important mediators of the disease. These experiments will lead to better treatment of the disease progression by affording the possibility of

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combinational therapies (e.g., multiple ribozymes targeted to different genes, ribozymes coupled with known small molecule inhibitors, or intermittent treatment with combinations of ribozymes and/or other chemical 5 biological molecules).

Other features and advantages of the invention will be apparent from the following description of the preferred embodiments thereof, and from the claims.

Description of the Preferred Embodiments

The drawings will first briefly be described.

Drawings

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Fig. 1 is a diagrammatic representation of a hammerhead motif ribozyme showing stems I, II and III (marked (I), (II) and (III) respectively) interacting with 15 a target region. The 5' and 3' ends of both ribozyme and target are Dashes indicate shown. base-paired nucleotides.

Figs. 2 - 11 are preferred targets for mdr-1 gene. chronic myelogenous leukemia, promyelocytic 20 leukemia, lymphoma Burkitt's lymphocytic or acute leukemia, follicular lymphoma, B-cell acute lymphocytic leukemia, breast cancer, colon carcinoma, neuroblastoma, and lung cancer, respectively. Target Sites

Ribozymes targeting selected regions of mRNA associated with tumor cell growth are preferably chosen to cleave the target RNA in a manner which inhibits translation of the mRNA. Genes are selected such that inhibition of translation will preferably inhibit cell 30 replication, <u>e.g.</u>, by inhibiting production of a necessary protein. Selection of effective target sites within these critical regions of mRNA entails testing the accessibility the target mRNA to hybridization with various oligonucleotide probes. These studies can be performed 35 using RNA or DNA probes and assaying accessibility by cleaving the hybrid molecule with RNaseH (see below). Alternatively, such a study can use ribozyme probes

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designed from secondary structure predictions of mRNAs, and assaying cleavage products by polyacrylamide gel electrophoresis (PAGE), to detect the presence of cleaved and uncleaved molecules.

The following are examples of cancer conditions which can be targetted in this manner. Chronic Myelogenous Leukemia

leukemia exhibits myelogenous Chronic characteristic disease course, presenting initially as a chronic granulocytic hyperplasia, and invariably evolving into an acute leukemia which is caused by the clonal expansion of a cell with a less differentiated phenotype (i.e., the blast crisis stage of the disease). CML is an unstable disease which ultimately progresses to a terminal stage which resembles acute leukemia. This lethal disease approximately 16,000 patients a affects Chemotherapeutic agents such as hydroxyurea or busulfan can reduce the leukemic burden but do not impact the life expectancy of the patient (e.g., approximately 4 years). 20 Consequently, CML patients are candidates for bone marrow However, for those transplantation (BMT) therapy. patients which survive BMT, disease recurrence remains a major obstacle. Apperley et al., 69 Br. J. Haematol. 239, 1988.

The Philadelphia (Ph) chromosome which results from the translocation of the abl oncogene from chromosome 9 to the bcr gene on chromosome 22 is found in greater than 95% of CML patients and in 10-25% of all cases of acute lymphoblastic leukemia (ALL). Fourth International 30 Workshop on Chromosomes in Leukemia. 11 Cancer Genet. Cytogenet. 316, 1982. In virtually all Ph-positive CMLs and approximately 50% of the Ph-positive ALLs, the leukemic cells express bcr-abl fusion mRNAs in which exon 2 (b2a2 junction) or exon 3 (b3a2 junction) from the major 35 breakpoint cluster region of the bcr gene is spliced to exon 2 of the abl gene. Heisterkamp et al., 315 Nature 758, 1985, Shtivelman et al., 69 <u>Blood</u> 971, 1987. In the

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remaining cases of Ph-positive ALL, the first exon of the bcr gene is spliced to exon 2 of the abl gene. Hooberman et al., 86 Proc. Natl. Acad. Sci. USA 4259, 1989, Heisterkamp et al., 16 Nucleic Acids Research 10069, 1988.

The b3a2 and b2a2 fusion mRNAs encode 210 kd bcr-abl fusion proteins which exhibit oncogenic activity. Daley et al., 247 Science 824, 1990, Heisterkamp et al., 344 Nature 251, 1990. The importance of the bcr-abl fusion protein (p210bcr-abl) in the evolution and maintenance of the leukemic phenotype in human disease has been demonstrated using antisense oligonucleotide inhibition of p210bcr-abl expression. These inhibitory molecules have been shown to inhibit the in vitro proliferation of leukemic cells in bone marrow from CML patients. Szczylik et al., 15 253 <u>Science</u> 562, 1991.

c-Myc Gene

c-Myc, when activated, can induce malignancy in a variety of tissues, most notably hematopoietic tissues (Leder et al., 222 Science 765, 1983). The most common 20 mechanism of c-myc activation is translocation to any of the immunoglobulin (Ig) or T-cell receptor loci during lymphoid maturation (Croce and Nowell, 65 Blood 1, 1985; Klein and Klein, 6 Immunol. Today 208, 1985). example, in Burkitt's lymphoma the c-myc locus on 25 chromosome 8 translocates most often to the Ig heavy chain locus on chromosome 14, but also to the lambda or kappa light chain Ig genes on chromosomes 2 and 22 (Magrath, in "Epstein-Barr Virus and Associated Diseases", M. Nijhoff Publishing: 631, 1986). In some instances the c-myc transcription region is altered in the non-coding exon 1 region; in such cases transcription is initiated at a cryptic promoter present in the first intron of the c-myc These rearrangements are thought to lead to locus. deregulation of c-myc expression.

c-Myc is not normally expressed in quiescent cells, but is temporally expressed in actively-dividing cells, most prominently during transition from Go to G1 phases of growth induction.

Experiments with transfected cell lines and transgenic animals have shown that c-myc activation plays a critical role, but is not sufficient for transformation (Adams et al., 318 Nature 533, 1985; Lombardi et al., 49 Cell 161, 1987; Schwartz et al., 6 Mol. Cell. Biol. 3221, 1986; Langdon et al., 47 Cell 11, 1986). Targeted inhibition of c-myc expression in tumor cell lines using antisense oligonucleotides has shown that c-myc expression is required for growth in certain lymphomas (McManaway et al., 335 Lancet 808, 1990).

Bcl-2 Gene

The bcl-2 gene is abnormally expressed in about 15 85% of follicular lymphomas and about 20% of diffuse t(14;18)(q32;q21) chromosomal a lymphomas due to rearrangement between the bcI-2 locus on chromosome 18 and the immunoglobulin heavy chain locus on chromosome 14 This (Yunis et al., 316 N. Engl. J. Med. 79, 1987). 20 chromosomal rearrangement represents the most common found in lymphoid malignancies in humans. A bcl-2/IgH fusion message is expressed; however, the bcl-2 protein-coding region is not interrupted since the major breakpoint region lies in the 3' nontranslated region of the bc1-2 25 transcript (Cleary et al., 47 <u>Cell</u> 19, 1986). gene represents a new form of proto-oncogene in that it encodes a mitochondrial protein which inhibits cell senescence (Hockenbery et al., 348 Nature 334, 1990), leading to extended survival of B-cells transfected with 30 this gene (Nunez et al., 86 Proc. Natl. Acad. Sci. USA 4589, 1989).

At least three different forms of bcl-2 mRNAs are found in pre-B-cells and T-cells, which vary due to alternative splicing and promoter usage. Two different proteins are produced, a 21 kD and a 26 kD peptide which vary at their carboxytermini. Both forms have identical

N termini encoded in exon 2 of the gene. Consequently, this region would be suitable for ribozyme targeting.

Breast Cancer

The epidermal growth factor (EGF) receptors have 5 been implicated in human cancer more frequently than any other family of growth factor receptors. The EGF receptor gene is often amplified or overexpressed in squamous cell carcinomas and glioblastomas. Jenkins et al., 39 Cancer Genet. Cytogenet. 253, 1989. Similarly, erbB-2 is often 10 overexpressed in adenocarcinomas of the stomach, breast Turc-Carel et al., 12 ibid. 1, and ovary. under Overexpression of either gene experimental conditions confers the transformed phenotype. Heim et al., 32 <u>ibid</u>. 13, 1988. In certain breast 15 carcinomas, the erbB-3 gene is overexpressed. Boehm et al., 7 EMBO J. 385, 1988.

The high incidence of human breast cancer has prompted efforts to model the disease in transgenic mice. The myc gene is amplified in some human breast cancers, 20 Escot et al., 83 Proc. Natl. Acad. Sci. USA 4834, 1986, and ras mutations have been observed. Barbacid, 56 Ann. Rev. Biochem. 779, 1987. Reproduction of disease by expression of the myc or ras genes in mice have given only sporadic results.

Breast cancer progression often correlates with amplification of the tyrosine kinase receptor gene denoted as c-erb-B2 or neu. The ligand for this receptor is unknown. Male and female mice expressing the neu gene both synchronously developed adenocarcinomas encompassing the entire gland. Muller et al., 54 <u>Cell</u> 105, 1988. Other strains developed tumors stochastically. Bouchard et al., 57 <u>ibid</u>. 931, 1989.

Colon Carcinoma

The platelet derived growth factor (PDGF) system

35 has served as a prototype for identification of substrates
of the receptor tyrosine kinases. Certain enzymes become
activated by the PDGF receptor kinase, including

phospholipase C and phosphatidylinositol 3' kinase, Ras guanosine triphosphate (GTPase) activating protein (GAP) and src-like tyrosine kinases. GAP regulates the function of the Ras protein. It stimulates the GTPase activity of the 21 kD Ras protein. Barbacid, 56 Ann. Rev. Biochem. 779, 1987. Microinjection of oncogenically activated Ras into NIH 3T3 cells induces DNA synthesis. Mutations that cause oncogenic activation of ras lead to accumulation of Ras bound to GTP, the active form of the molecule. These mutations block the ability of GAP to convert Ras to the inactive form. Mutations that impair the interactions of Ras with GAP also block the biological function of Ras.

While a number of ras alleles exist (N-ras, K-ras, H-ras) which have been implicated in carcinogenesis, the type most often associated with colon and pancreatic carcinomas is the K-ras. Ribozymes which are targeted to certain regions of the K-ras allelic mRNAs may also prove inhibitory to the function of the other allelic mRNAs of the N-ras and H-ras genes.

20 Lung Cancer/L-myc Gene

Expression of the myc oncogene is known to alter cell growth in a number of tissues. The product of this gene is a protein which is known to be a transcriptional activator that can act singly or in combination with other oncogene proteins. The L-myc gene is often activated by translocations of DNA from other regions of the genome to the regulatory regions 5' of the myc gene ORF. After transcription of the L-myc mRNA, alternate splicing of the transcript is known to occur. Kaye et al., 8 Mol. Cell Biol. 196, 1988. The alternate mRNAs produced contain a common 5' exon 1 and portions of a common exon 2. These common regions of mRNA structure allow the use of nucleic acid targeted therapeutics which can inactivate both species of mRNA with one therapeutic molecule.

35 Promyelocytic Leukemia

Acute promyelocytic leukemia is characterized by a specific translocation, a t(15;17)(q22;q11.2-12), which

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is found in some 90% of the cases. The t(15;17) is often the only detectable cytogenetic abnormality present in the leukemic cells. This rearrangement results in the fusion of two genes, the promyelocytic leukemia gene (PML) on 5 chromosome 15, and the retinoic acid receptor alpha gene (RARA) on chromosome 17 (J. Borrow et al., 249 Science 1577, 1990; H. de Thé et al., 347 Nature 558, 1990). RARA is a hormonally-responsive transcriptional regulatory protein, while the function of the PML is as yet unknown.

A fusion message is expressed in the leukemic cells which encodes the N-terminal coding region of the PML gene and the C-terminal coding region of the RARA gene. Expression of this fusion gene apparently inhibits normal myeloid differentiation. The biological relevance 15 of this rearrangement to the etiology of the disease has been exemplified by the discovery that all-trans retinoic acid can be used to achieve complete clinical remission, presumably by inducing differentiation of the leukemic This suggests that the fusion protein is still cells. 20 hormonally responsive.

The treatment of leukemic cells with retinoic acid is not preferable over the long term because retinoic acid is a generalized inducer of differentiation in all Thus, systemic cell types, not just leukemic cells. 25 administration of these compounds can lead to a number of deleterious side effects by differentiating cells which should not be in a differentiated state. A treatment which gives suppression of the transformed phenotype in leukemic cells without affecting other cell types is 30 preferable, as described herein.

B-Cell Acute Lymphocytic Leukemia

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Leukemia comprises some 3% of the new cancer cases per year, with lymphocytic leukemias accounting for approximately half (National Cancer Institute, 1990 statistics). A subset of lymphocytic leukemias of the acute pre-B-cell type are associated with a specific chromosomal translocation, a t(1;19)(q23;p13.3)

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Kamps et al., 60 <u>Cell</u> 547, 1990; J. Nourse et al., <u>ibid</u>. p.535). This rearrangement results in the fusion of two genes, the PBX gene present on chromosome 1, and the E2A gene present on chromosome 19. While the E2A transcript 5 is found in all B-cell types, the PRL gene is not normally expressed in B-cells. However, an E2A/PRL fusion message is constitutively expressed from this aberrant locus in the leukemic cells. This fusion message encodes the Nterminal region of the E2A, including the transcriptional 10 activating domain of that gene, and the C-terminal region of the PRL gene, which contains a homeodomain DNA binding a potentially functional chimeric Thus, transcriptional regulatory protein is expressed in the leukemic cells.

The PRL sequences found in the fusion mRNA are good targets for ribozyme therapy since PRL is not expressed in non-leukemic B-cells. Whether the E2A sequences can be targeted by ribozymes is unclear since such ribozymes may inhibit E2A expression in normal Bcells. It is not known how normal B-cells are affected by 20 inhibition of E2A.

The following is but one example of a method by which suitable target sites can be identified and is not Generally, the method limiting in this invention. involves identifying potential cleavage sites for a hammerhead ribozyme, and then testing each of these sites to determine their suitability as targets by ensuring that secondary structure formation is minimal.

are compared The mRNA sequences Putative ribozyme cleavage appropriate target region. sites are identified from weak or non-base paired regions of the mRNA. These sites represent the preferred sites for hammerhead or other ribozyme cleavage within these target mRNAs.

Short RNA substrates corresponding to each of the mRNA sites are designed. Each substrate is composed of two to three nucleotides at the 5' and 3' ends that

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will not base pair with a corresponding ribozyme recognition region. The unpaired regions flank a central region of 12-14 nucleotides to which complementary arms in the ribozyme are designed.

The structure of each substrate sequence is predicted using a PC fold computer program. Sequences which give a positive free energy of binding are accepted. Sequences which give a negative free energy are modified by trimming one or two bases from each of the ends. If the modified sequences are still predicted to have a strong secondary structure, they are rejected.

After substrates are chosen, ribozymes are designed to each of the RNA substrates. Ribozyme folding is also analyzed using PC fold.

Ribozyme molecules are sought which form hammerhead motif stem II (see Fig. 1) regions and contain flanking arms which are devoid of intramolecular base pairing. Often the ribozymes are modified by trimming a base from the ends of the ribozyme, or by introducing additional base pairs in stem II to achieve the desired fold. Ribozymes with incorrect folding are rejected. After substrate/ribozyme pairs are found to contain correct intramolecular structures, the molecules are folded together to predict intermolecular interactions.

25 A schematic representation of a ribozyme with its coordinate base pairing to its cognate target sequence is shown in Fig. 1. Examples of useful targets are listed in Figs. 2 - 11.

Those targets thought to be useful as ribozyme 30 targets can be tested to determine accessibility to nucleic acid probes in a ribonuclease H assay (see below). This assay provides a quick test of the use of the target site without requiring synthesis of a ribozyme. It can be used to screen for sites most suited for ribozyme attack.

35 Synthesis of Ribozymes

Ribozymes useful in this invention can be produced by gene transcription as described by Cech,

supra, or by chemical synthesis. Chemical synthesis of RNA is similar to that for DNA synthesis. The additional 2'-OH group in RNA, however, requires a different protecting group strategy to deal with selective 3'-5' internucleotide bond formation, and with RNA susceptibility to degradation in the presence of bases. The recently developed method of RNA synthesis utilizing the t-butyldimethylsilyl group for the protection of the 2' hydroxyl is the most reliable method for synthesis of ribozymes. The method reproducibly yields RNA with the correct 3'-5' internucleotide linkages, with average coupling yields in excess of 99%, and requires only a two-step deprotection of the polymer.

A method based upon H-phosphonate chemistry of 15 phosphoramidites gives a relatively lower efficiency than a method based upon phosphoroamidite This is a problem for synthesis of DNA as chemistry. A promising approach to scale-up of automatic oligonucleotide synthesis has been described recently for 20 the H-phosphonates. A combination of a proper coupling time and additional capping of "failure" sequences gave high yields in the synthesis of oligodeoxynucleotides in scales in the range of 14 μ moles with as little as 2 equivalents of a monomer in the coupling step. 25 alternative approach is to use soluble polymeric supports (e.g., polyethylene glycols), instead of the conventional can vield This method solid supports. oligonucleotides in hundred milligram quantities per batch utilizing about 3 equivalents of a monomer in a coupling 30 step.

Various modifications to ribozyme structure can be made to enhance the utility of ribozymes. Such modifications will enhance shelf-life, half-life in vitro, stability, and ease of introduction of such ribozymes to the target site, <u>e.g.</u>, to enhance penetration of cellular membranes, and confer the ability to recognize and bind to targeted cells.

Exogenous delivery of ribozymes benefits from chemical modification of the backbone, e.g., by the overall negative charge of the ribozyme molecule being reduced to facilitate diffusion across the cell membrane. 5 The present strategies for reducing the oligonucleotide charge include: modification of internucleotide linkages by methylphosphonates, use of phosphoramidites, linking oligonucleotides to positively charged molecules, creating complex packages composed of oligonucleotides, 10 lipids and specific receptors or effectors for targeted Examples of such modifications include sulfurcontaining ribozymes containing phosphorothicates and phosphorodithioates as internucleotide linkages in RNA. Synthesis of such sulfur-modified ribozymes is achieved by 15 use of the sulfur-transfer reagent, 3H-1,2-benzenedithiol-3-one 1,1-dioxide. Ribozymes may also contain ribose Pyrimidine analogues are modified ribonucleotides. prepared from uridine using a procedure employing diethylamino sulphur trifluoride (DAST) as a starting 20 reagent. Ribozymes can also be either electrostatically or covalently attached to polymeric cations for the purpose of reducing charge. The polymer can be attached to the ribozyme by simply converting the 3'-end to a ribonucleoside dialdehyde which is obtained by a periodate 25 cleavage of the terminal 2',3'-cis diol system. Depending on the specific requirements for delivery systems, other possible modifications may include different linker arms containing carboxyl, amino or thiol functionalities. Yet further examples include use of methylphosphonates and 2'-30 O-methylribose and 5' or 3' capping or blocking with m₇GpppG or m₃^{2,2,7}GpppG.

For example, a kinased ribozyme is contacted with guanosine triphosphate and guanyltransferase to add a m³G cap to the ribozyme. After such synthesis, the ribozyme can be gel purified using standard procedure. To ensure that the ribozyme has the desired activity, it may be tested with and without the 5' cap using standard

procedures to assay both its enzymatic activity and its stability.

Synthetic ribozymes, including those containing various modifiers, can be purified by high pressure liquid Other liquid chromatography (HPLC). 5 chromatography techniques, employing reverse phase columns and anion exchangers on silica and polymeric supports may also be useđ.

There follows an example of the synthesis of one 10 ribozyme. A solid phase phosphoramidite chemistry was employed. Monomers used were 2'-tert-butyl-dimethylsilyl cyanoethylphosphoramidities of uridine, cytosine, N-phenoxyacetyl adenosine and guanosine (Glen Solid phase synthesis was Research, Sterling, VA). 15 carried out on either an ABI 394 or 380B DNA/RNA synthesizer using the standard protocol provided with each machine. The only exception was that the coupling step was increased from 10 to 12 minutes. The phosphoramidite concentration was 0.1 M. Synthesis was done on a 1 μ mole 20 scale using a 1 μ mole RNA reaction column (Glen Research). The average coupling efficiencies were between 97% and 98% for the 394 model, and between 97% and 99% for the 380B model, as determined by a calorimetric measurement of the released trityl cation.

Blocked ribozymes were cleaved from the solid support (e.g., CPG), and the bases and diphosphoester moiety deprotected in a sterile vial by dry ethanolic ammonia (2 mL) at 55°C for 16 hours. The reaction mixture was cooled on dry ice. Later, the cold liquid was 30 transferred into a sterile screw cap vial and lyophilized.

To remove the 2'-tert-butyl-dimethylsilyl groups from the ribozyme, the residue was suspended in 1 M tetran-butylammonium fluoride in dry THF (TBAF), using a 20fold excess of the reagent for every silyl group, for 16 35 hours at ambient temperature (about 15-25°C). reaction was quenched by adding an equal volume of sterile WO 93/23057 PCT/US93/04573

1 M triethylamine acetate, pH 6.5. The sample was cooled and concentrated on a SpeedVac to half the initial volume.

The ribozymes were purified in two steps by HPLC on a C4 300 Å 5 mm DeltaPak column in an acetonitrile 5 gradient.

The first step, or "trityl on" step, was a separation of 5'-DMT-protected ribozyme(s) from failure sequences lacking a 5'-DMT group. Solvents used for this step were: A (0.1 M triethylammonium acetate, pH 6.8) and B (acetonitrile). The elution profile was: 20% B for 10 minutes, followed by a linear gradient of 20% B to 50% B over 50 minutes, 50% B for 10 minutes, a linear gradient of 50% B to 100% B over 10 minutes, and a linear gradient of 100% B to 0% B over 10 minutes.

The 'second step was a purification of a completely deblocked ribozyme by a treatment of 2% trifluoroacetic acid on a C4 300 Å 5 mm DeltaPak column in an acetonitrile gradient. Solvents used for this second step were: A (0.1 M triethylammonium acetate, pH 6.8) and B (80% acetonitrile, 0.1 M triethylammonium acetate, pH 6.8). The elution profile was: 5% B for 5 minutes, a linear gradient of 5% B to 15% B over 60 minutes, 15% B for 10 minutes, and a linear gradient of 15% B to 0% B over 10 minutes.

25 The fraction containing ribozyme was cooled and lyophilized on a SpeedVac. Solid residue was dissolved in a minimum amount of ethanol and sodium perchlorate in acetone. The ribozyme was collected by centrifugation, washed three times with acetone, and lyophilized.

30 Expression Vector

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While synthetic ribozymes are preferred in this invention, those produced by expression vectors can also be used. In designing a suitable ribozyme expression vector the following factors are important to consider. The final ribozyme must be kept as small as possible to minimize unwanted secondary structure within the ribozyme.

A promoter (e.g., a T7, human cytomegalovirus immediate

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early (iel), human beta actin, or U6 snRNA promoters) should be chosen to be a relatively strong promoter, and expressible both in vitro and in vivo (e.g., by coinfection with the T7 RNA polymerase gene, the human 5 cytomegalovirus immediate early (iel) or human beta actin Such a promoter should express the ribozyme promoters). at a level suitable to effect production of enough ribozyme to destroy a target RNA, but not at too high a level to prevent other cellular activities from occurring (unless cell death itself is desired).

A hairpin at the 5' end of the ribozyme is ensure that the required transcription useful initiation sequence (GG or GGG or GGGAG) does not bind to some other part of the ribozyme and thus affect regulation The 5' hairpin is also 15 of the transcription process. useful to protect the ribozyme from 5'-3' exonucleases. A selected hairpin at the 3' end of the ribozyme gene is useful since it acts as a transcription termination signal, and protects the ribozyme from 3'-5' exonuclease One example of a known termination signal is activity. that present on the T7 RNA polymerase system. This signal is about 30 nucleotides in length. Other 3' hairpins of shorter length can be used to provide good termination and RNA stability. Such hairpins can be inserted within the 25 vector sequences to allow standard ribozymes to be placed in an appropriate orientation and expressed with such sequences attached.

Poly(A) tails are also useful to protect the 3' end of the ribozyme. These can be provided by either including a poly(A) signal site in the expression vector (to signal a cell to add the poly(A) tail in vivo), or by introducing a poly(A) sequence directly expression vector. In the first approach the signal must located to prevent unwanted secondary formation with other parts of the ribozyme. In the second approach, the poly(A) stretch may reduce in size over time when expressed in vivo, and thus the vector may need to be

checked over time. Care must be taken in addition of a poly(A) tail which binds poly(A) binding proteins which prevent the ribozyme from acting.

Ribozyme Testing

Once the desired ribozymes are selected, synthesized and purified, they are tested in kinetic and other experiments to determine their utility. An example of such a procedure is provided below.

Preparation of Ribozyme

Crude synthetic ribozyme (typically 350 µg at a time) was purified by separation on a 15% denaturing polyacrylamide gel (0.75 mm thick, 40 cm long) and Once excised, gel slices visualized by UV shadowing. containing full length ribozyme were soaked in 5 ml gel elution buffer (0.5 M NH,OAc, 1 mM EDTA) overnight with shaking at 4°C. The eluent was desalted over a C-18 matrix (Sep-Pak cartridges, Millipore, Milford, MA) and vacuum dried. The dried RNA was resuspended in $50-100~\mu l$ TE (TRIS 10 mM, EDTA 1 mM, pH 7.2). An aliquot of this 20 solution was diluted 100-fold into 1 ml TE, half of which was used to spectrophotometrically quantitate the ribozyme The concentration of this dilute stock was solution. Purity of the ribozyme was typically 150-800 nM. confirmed by the presence of a single band on a denaturing 25 polyacrylamide gel.

A ribozyme may advantageously be synthesized in two or more portions. Each portion of a ribozyme will generally have only limited or no enzymatic activity, and the activity will increase substantially (by at least 5-10 30 fold) when all portions are ligated (or otherwise juxtaposed) together. A specific example of hammerhead ribozyme synthesis is provided below.

The method involves synthesis of two (or more) shorter "half" ribozymes and ligation of them together using T4 RNA ligase. For example, to make a 34 mer one synthesized, 17 mers are ribozyme, phosphorylated, and both are gel purified. These purified 17 mers are then annealed to a DNA splint strand complementary to the two 17 mers. (Such a DNA splint is not always necessary.) This DNA splint has a sequence designed to locate the two 17 mer portions with one end of each adjacent each other. The juxtaposed RNA molecules are then treated with T4 RNA ligase in the presence of ATP. The 34 mer RNA so formed is then HPLC purified.

Preparation of Substrates

Approximately 10-30 pmoles of unpurified substrate was radioactively 5' end-labeled with T4 polynucleotide kinase using 25 pmoles of $[\gamma^{-32}P]$ ATP. The entire labeling mix was separated on a 20% denaturing polyacrylamide gel and visualized by autoradiography. The full length band was excised and soaked overnight at 4°C in 100 μ l of TE (10 mM Tris-HCl pH 7.6, 0.1 mM EDTA).

Kinetic Reactions

For reactions using short substrates (between 8 and 16 bases) a substrate solution was made 1X in assay buffer (75 mM Tris-HCl, pH 7.6; 0.1 mM EDTA, 10 mM MgCl₂) such that the concentration of substrate was less than 1 nM. A ribozyme solution (typically 20 nM) was made 1X in assay buffer and four dilutions were made using 1X assay buffer. Fifteen µl of each ribozyme dilution (i.e., 20, 16, 12, 8 and 4 nM) was placed in a separate tube. These tubes and the substrate tube were pre-incubated at 37°C for at least five minutes.

The reaction was started by mixing 15 µl of substrate into each ribozyme tube by rapid pipetting (note that final ribozyme concentrations were 10, 8, 6, 4, 30 2 nM). Five µl aliquots were removed at 15 or 30 second intervals and quenched with 5 µl stop solution (95% formamide, 20 mM EDTA xylene cyanol, and bromphenol blue dyes). Following the final ribozyme time point, an aliquot of the remaining substrate was removed as a zero ribozyme control.

The samples were separated on either 15% or 20% polyacrylamide gels. Each gel was visualized and

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quantitated with an Ambis beta scanner (Ambis Systems, San Diego, CA).

For the most active ribozymes, kinetic analyses were performed in substrate excess to determine K_m and K_{cat} 5 values.

For kinetic reactions with long RNA substrates (greater than 15 bases in length) the substrates were prepared by transcription using T7 RNA polymerase and defined templates containing a T7 promoter, and DNA encoding appropriate nucleotides of the target RNA. The substrate solution was made 1X in assay buffer (75 mM Tris-HCl, pH 7.6; 0.1 mM EDTA; 10 mM MgCl₂) and contained 58 nanomolar concentration of the long RNA molecules. The reaction was started by addition of gel purified ribozymes to 1 μM concentration. Aliquots were removed at 20, 40, 60, 80 and 100 minutes, then quenched by the addition of 5 μl stop solution. Cleavage products were separated using denaturing PAGE. The bands were visualized and quantitated with an Ambis beta scanner.

20 <u>Kinetic Analysis</u>

A simple reaction mechanism for ribozyme-mediated cleavage is:

$$k_1 k_2$$

$$R + S = [R:S] = [R:P] \longrightarrow R + P$$

$$k_{-1}$$

where R = ribozyme, S = substrate, and P = products. The boxed step is important only in substrate excess. Because 30 ribozyme concentration is in excess over substrate concentration, the concentration of the ribozyme-substrate complex ([R:S]) is constant over time except during the very brief time when the complex is being initially formed, i.e.,:

$$\frac{d[R:S]}{dt} = 0$$

where t = time, and thus:

$$(R)(S)k_1 = (RS)(k_2 + k_1).$$

The rate of the reaction is the rate of disappearance of substrate with time:

Rate = $\frac{-d(S)}{dt}$ = $k_2(RS)$

5 Substituting these expressions:

(R) (S)
$$k_1 = 1/k_2 -d(S) (k_2 + k_1)$$

or:

 $\frac{-d(S)}{S} = \frac{k_1 k_2}{(k_2 + k_1)}$ (R) dt

Integrating this expression with respect to time yields:

$$-\ln \frac{S}{S_0} = \frac{k_1 k_2}{(k_2 + k_1)}$$
 (R) t

where S_0 = initial substrate. Therefore, a plot of the 15 negative log of fraction substrate uncut versus time (in minutes) yields a straight line with slope:

slope =
$$\frac{k_1 k_2}{(k_2 + k_1)}$$
 (R) = k_{obs}

where k_{obs} = observed rate constant. A plot of slope (k_{obs}) 20 versus ribozyme concentration yields a straight line with a slope which is:

slope =
$$\frac{k_1k_2}{(k_2 + k_1)}$$
 which is $\frac{k_{\text{cat}}}{K_m}$

Using these equations the data obtained from the 25 kinetic experiments provides the necessary information to determine which ribozyme tested is most useful, or active. Such ribozymes can be selected and tested in *in vivo* or ex vivo systems.

Liposome Preparation

organic solvent (CHCl₃, methanol, diethylether, ethanol, etc.). The organic solvent is removed by evaporation. The lipid is hydrated into suspension with 0.1x phosphate buffered saline (PBS), then freeze-thawed 3x using liquid nitrogen and incubation at room temperature. The

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suspension is extruded sequentially through a 0.4 μm , $0.2~\mu\text{m}$ and $0.1~\mu\text{m}$ polycarbonate filters at maximum pressure of 800 psi. The ribozyme is mixed with the extruded liposome suspension and lyophilized to dryness. 5 The lipid/ribozyme powder is rehydrated with water to onetenth the original volume. The suspension is diluted to the minimum volume required for extrusion (0.4 ml for 1.5 ml barrel and 1.5 ml for 10 ml barrel) with 1xPBS and re-extruded through 0.4 μm, 0.2 μm, 0.1 μm polycarbonate The liposome entrapped ribozyme is separated 10 filters. from untrapped ribozyme by gel filtration chromatography (SEPHAROSE CL-4B, BIOGEL A5M). The liposome extractions are pooled and sterilized by filtration through a 0.2 μm The free ribozyme is pooled and recovered by filter. The liposome concentration is 15 ethanol precipitation. determined by incorporation of a radioactive lipid. ribozyme concentration is determined by labeling with 32P. Rossi et al., 1992, supra (and references cited therein) describe other methods suitable for preparation of liposomes. 20

Examples of other useful liposome preparations which display similar degrees of uptake of both a radioactive lipid marker and an entrapped fluorophore by Vero cells showed different fluorescent staining patterns. 25 Specifically, liposomes composed of DPPG/DPPC/Cholesterol 50/17/33) gave a punctate pattern of (in a ratio of: fluorescence, while DOPE/Egg PC/Cholesterol (30/37/33) gave a diffuse, homogeneous pattern of fluorescence in the Cell fractionation showed that 80% of the cytoplasm. DPPG/DPPC/Cholesterol from the 30 entrapped contents formulation was localized in the membrane fraction, whereas the DOPE/Egg PC/Cholesterol formulation was localized in the cytoplasm. Further characterization of the latter formulation showed that after 3 hours, 70% of the fluorescence was cytoplasmic and 30% was in the 35 membrane. After 24 hours, uptake had increased 5-fold and

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the liposome contents were distributed 50/50 between the cytoplasmic and membrane fractions.

Liposomes containing 15 ribozymes (32P-labeled) targeted to the HSV ICP4 mRNA were prepared and incubated 5 with the cells. After 24 hours, 25% of the liposome dose was taken up with approximately 60,000 liposomes per cell. Thirty percent of the delivered ribozyme was intact after 24 hours. Cell fractionation studies showed 40% of the intact ribozyme to be in the membrane fraction and 52% of the intact ribozyme to be in the cytoplasmic fraction.

<u>In Vivo Assay</u>

The efficacy of action of a chosen ribozyme may be tested in vivo by use of cell cultures sensitive to mdr-1 gene expression, using standard procedures in transformed cells or animals which express the target mRNA using standard procedures.

The efficacy of action of a chosen ribozyme may be tested in tissue culture by use of transformed cells containing the target mRNA (e.g., K562 cells which express the b3a2 fusion mRNA) using standard procedures. Alternatively, ribozyme efficacy could be tested with peripheral blood or bone marrow from CML patients using soft-agar colony forming assays. Such methods are known to those educated in this field.

Ribonuclease Protection Assay

The accumulation of target mRNA in cells or the cleavage of the mRNA by ribozymes or RNaseH (in vitro or in vivo) can be quantified using an RNase protection assay.

In this method, antisense riboprobes are transcribed from template DNA using T7 RNA polymerase (U.S. Biochemical) in 20 µl reactions containing 1X transcription buffer (supplied by the manufacturer), 0.2 mM ATP, GTP and UTP, 1 U/µl pancreatic RNase inhibitor (Boehringer Mannheim Biochemicals) and 200 µCi ³²P-labeled CTP (800 Ci/mmol, New England Nuclear) for 1 hour at 37°C. Template DNA is digested with 1 U RNase-free DNaseI (U.S.

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Biochemical, Cleveland, OH) at 37°C for 15 minutes and unincorporated nucleotides removed by G-50 SEPHADEX spin chromatography.

In a manner similar to the transcription of antisense probe, the target mRNA can be transcribed in vitro using a suitable DNA template. The transcript is purified by standard methods and digested with ribozyme at 37°C according to methods described later.

Alternatively, afflicted (mRNA-expressing) cells 10 expressing the target mRNA bcr-abl fusion transcript are harvested into 1 ml of PBS, transferred to a 1.5 ml EPPENDORF tube, pelleted for 30 seconds at low speed in a microcentrifuge, and lysed in 70 μ l of hybridization buffer (4 M quanidine isothiocyanate, 0.1% sarcosyl, 25 mM sodium citrate, pH 7.5). Cell lysate (45 μ l) or defined 15 amounts of in vitro transcript (also in hybridization buffer) is then combined with 5 μ l of hybridization buffer containing 5 x 10^5 cpm of each antisense riboprobe in 0.5 ml EPPENDORF tubes, overlaid with 25 µl mineral oil, and hybridization accomplished by heating overnight at 55°C. The hybridization reactions are diluted into 0.5 ml RNase solution (20 U/ml RNaseA, 2 U/ml RNaseT1, 10 U/ml RNasefree DNaseI in 0.4 M NaCl), heated for 30 minutes at 37°C, and 10 µl of 20% SDS and 10 µl of Proteinase K (10 mg/ml) 25 added, followed by an additional 30 minutes incubation at Hybrids are partially purified by extraction with 0.5 ml of a 1:1 mixture of phenol/chloroform; aqueous phases are combined with 0.5 ml isopropanol, and RNaseresistant hybrids pelleted for 10 minutes at room 30 temperature (about 20°C) in a microcentrifuge. are dissolved in 10 µl loading buffer (95% formamide, 1X TBE, 0.1% bromophenol blue, 0.1% xylene cylanol), heated to 95°C for five minutes, cooled on ice, and analyzed on under denaturing polyacrylamide/7 M urea gels 35 conditions.

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Ribozyme Stability

The chosen ribozyme can be tested to determine its stability, and thus its potential utility. Such a test can also be used to determine the effect of various chemical modifications (e.g., addition of a poly(A) tail) on the ribozyme stability and thus aid selection of a more stable ribozyme. For example, a reaction mixture contains 1 to 5 pmoles of 5' (kinased) and/or 3' labeled ribozyme, 15 μg of cytosolic extract and 2.5 mM MgCl₂ in a total volume of 100 μl. The reaction is incubated at 37°C. Eight μl aliquots are taken at timed intervals and mixed with 8 μl of a stop mix (20 mM EDTA, 95% formamide). Samples are separated on a 15% acrylamide sequencing gel, exposed to film, and scanned with an Ambis.

A 3'-labeled ribozyme can be formed by incorporation of the $^{32}\text{P-labeled}$ cordycepin at the 3' OH using poly(A) polymerase. For example, the poly(A) polymerase reaction contains 40 mM Tris, pH 8, 10 mM MgCl₂, 250 mM NaCl, 2.5 mM MnCl₂; 3 μ l ^{32}P cordycepin, 500 Ci/mM; and 6 units poly(A) polymerase in a total volume of 50 μ l. The reaction mixture is incubated for 30 minutes at 37°C.

Effect of Base Substitution upon Ribozyme Activity

structural which determine primary To cleavage 25 characteristics could change ribozyme substrate, minor base changes can be made in the substrate cleavage region recognized by a specific ribozyme. example, the substrate sequences can be changed at the central "C" nucleotide, changing the cleavage site from a GUC to a GUA motif. The $K_{\text{cat}}/K_{\text{m}}$ values for cleavage using each substrate are then analyzed to determine if such a change increases ribozyme cleavage rates. Similar experiments can be performed to address the effects of changing bases complementary to the ribozyme binding arms. Changes predicted to maintain strong binding to the complementary substrate are preferred. Minor changes in alter ribozyme/substrate nucleotide content can

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interactions in ways which are unpredictable based upon binding strength alone. Structures in the catalytic core region of the ribozyme recognize trivial changes in either substrate structure or the three dimensional structure of the ribozyme/substrate complex.

To begin optimizing ribozyme design, the cleavage rates of ribozymes containing varied arm lengths, but targeted to the same length of short RNA substrate can be tested. Minimal arm lengths are required and effective cleavage varies with ribozyme/substrate combinations.

The cleavage activity of selected ribozymes can be assessed using target mRNA-homologous substrates. The assays are performed in ribozyme excess and approximate $K_{\rm cat}/K_{\rm min}$ values obtained. Comparison of values obtained with short and long substrates indicates utility in vivo of a ribozyme.

<u>Intracellular Stability of Liposome-Delivered</u> Ribozymes

To test the stability of a chosen ribozyme 20 in vivo the following test is useful. Ribozymes are 32Pend-labeled, entrapped in liposomes and delivered to target mRNA containing cells for three hours. The cells purified ribozyme fractionated and is are phenol/chloroform extraction. Alternatively, cells (1x10', 25 T-175 flask) are scraped from the surface of the flask are cultured and washed twice with cold PBS. The cells are homogenized by douncing 35 times in 4 ml of TSE (10 mM Tris, pH 7.4, 0.25 M Sucrose, mM EDTA). Nuclei are pelleted at 100xg for 10 minutes. Subcellular organelles (the membrane fraction) are pelleted at 200,000xg for two hours using an SW60 rotor. The pellet is resuspended in 1 ml of H buffer (0.25 M Sucrose, 50 mM HEPES, pH 7.4). The supernatant contains the cytoplasmic fraction (in approximately 3.7 ml). The nuclear pellet is resuspended in 1 ml of 65% sucrose in TM (50 mM Tris, pH 7.4, 2.5 mM MgCl₂) and banded on a sucrose step gradient (1 ml nuclei in 65% sucrose TM, 1 ml 60% sucrose TM, 1 ml 55% sucrose

TM, 50% sucrose TM, 300 µl 25% sucrose TM) for one hour at 37,000xg with an SW60 rotor. The nuclear band is harvested and diluted to 10% sucrose with TM buffer. Nuclei are pelleted at 37,000xg using an SW60 rotor for 15 minutes and the pellet resuspended in 1 ml of TM buffer. Aliquots are size fractionated on denaturing polyacrylamide gels and the intracellular localization determined. By comparison to the migration rate of newly synthesized ribozyme, the various fractions containing intact ribozyme can be determined.

To investigate modifications which would lengthen the half-life of ribozyme molecules intracellularly, the cells may be fractioned as above and the purity of each fraction assessed by assaying enzyme activity known to exist in that fraction.

The various cell fractions are frozen at -70°C and used to determine relative nuclease resistances of modified ribozyme molecules. Ribozyme molecules may be synthesized with 5 phosphorothicate (ps), or 2'-O-methyl 20 (2'-OMe) modifications at each end of the molecule. These molecules and a phosphodiester version of the ribozyme are end-labeled with 32P and ATP using T4 polynucleotide Equal concentrations are added to the cell kinase. cytoplasmic extracts and aliquots of each taken at 10 25 minute intervals. The samples are size fractionated by denaturing PAGE and relative rates of nuclease resistance analyzed by scanning the gel with an Ambis β -scanner. The results show whether the ribozymes are digested by the cytoplasmic extract, and which versions are relatively Modified ribozymes generally 30 more nuclease resistant. maintain 80-90% of the catalytic activity of the native ribozyme when short RNA substrates are employed.

Unlabeled, 5' end-labeled or 3' end-labeled ribozymes can be used in the assays. These experiments can also be performed with human cell extracts to verify the observations.

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In one example, Vero or HeLa cells were grown to 90-95% confluency in 175 cm2 tissue culture flasks, scraped into 10 ml of cold phosphate buffered saline (PBS), then washed once in 10 ml of cold PBS and once in 10 ml of cold 5 TSE (10 mM Tris, pH 7.4; 0.25 M sucrose; 1 mM EDTA). cell pellets were resuspended in 4 ml of TSE, dounced 35x on ice, and the released nuclei pelleted by centrifugation at 1000g for 10 minutes. The nuclear pellet was resuspended in 1 ml of 65% sucrose TM (50 mM Tris, pH 7.4; 10 2.5 mM MgCl₂) and transferred to Beckman ultra-clear tubes. The following sucrose TM solutions were layered on top of the sample: 1 ml 60%, 1 ml 55%, and 25% sucrose to the top of the tube. Gradients were spun in an SW60 rotor at 37,000g for 1 hour. HeLa nuclei banded at the 55-60% 15 sucrose boundary and Vero nuclei banded at the 50-55% sucrose boundary. Nuclear bands were harvested, diluted sucrose with TM buffer. and pelleted by centrifugation at 37,000g for 15 minutes using an SW60 rotor. The nuclear pellet was resuspended in 1 ml of TM 20 buffer. Subcellular organelles and membrane components in the post nuclear supernatant were separated from the cytoplasmic fraction by centrifugation at 200,000g for 2 hours in an SW60 rotor. The pellet contained the membrane fraction, which was resuspended in 1 ml of H buffer (0.25 25 M sucrose; 50 mM HEPES, pH 7.4), and the supernatant contained the cytoplasmic fraction.

Purity of the various fractions was assessed using enzymatic markers specific for the cytoplasmic and membranous fractions. Three enzyme markers for the 30 membranous fraction were used; hexosaminidase and β -glucocerebrosidase are localized in lysosomes, while alkaline phosphodiesterase is specific to endosomes. Specifically, the assays were as follows:

For N-acetyl-beta-hexosaminidase, the reaction mixture contained 0.3 mg/ml 4-methylumbelliferyl-N-acetyl-glucosaminide; 20 mM sodium citrate; pH 4.5; 0.01% Triton X-100; and 100 μ l of sample in a final volume of 500 μ l

(Harding et al., 64 Cell 393, 1991). The reactions were incubated at 37°C for 1 hour and stopped by the addition of 1.5 ml of stop buffer (0.13 M glycine, 0.07 M NaCl, 0.08 M sodium carbonate, pH 10.6). The reaction product fluorescence Hitachi F-4010 quantitated in a was spectrophotometer by excitation of the fluorophore at 360 nm and analysis of the emission at 448 nm.

For Alkaline Phosphodiesterase, the assay medium contained 25 mM CAPS (3-(Cyclohexylamino)-propanesulfonic 10 acid), pH 10.6; 0.05% Triton X-100; 15 mM MgCl₂; 1.25 mg/ml Thymidine-5'-monophosphate-p-nitrophenyl ester; and 100 μl of sample in a total reaction volume of 200 μ l. reactions were incubated at 37°C for 2 hours, then diluted to 1 ml with ${\rm H}_2{\rm O}$ and the absorbance was measured at 400 nm (Razell and Khorana, 234 J. Biol. Chem. 739, 1959).

For β -glucocerebrosidase, the reaction contained 85 nM sodium citrate, pH 5.9; 0.12% Triton X-100; 0.1% sodium taurocholate; 5 mM 4-methylumbelliferyl β -Dglucopyranoside; and 125 μl of sample in a total volume of 20 250 μl (Kennedy and Cooper, 252 Biochem. J. 739, 1988). The reaction was incubated at 37°C for 1 hour and stopped by the addition of 0.75 ml of stop buffer. formation was measured in a fluorescence spectrophotometer by using an excitation wavelength of 360 nm and analysis 25 of the emission at 448 nm.

marker, cytoplasmic enzyme The dehydrogenase, was assayed in an assay mixture containing 0.2 M Tris; pH 7.4; 0.22 mM NADH; 1 mM sodium pyruvate; and 50 μ l of sample in a final volume of 1.05 ml. Enzyme levels were determined by decreased absorbency at 350 nm resulting from the oxidation of NADH at room temperature (Silverstein and Boyer, 239 J. Biol. Chem. 3901, 1964).

Lactate dehydrogenase was found predominantly in the cytoplasmic fractions of both Vero and HeLa cells, while β -glucocerebrosidase and alkaline phosphodiesterase were found almost exclusively in the membranous fractions. The hexosaminidase activity in Vero cell fractions was concentrated in the membranous fraction (70%) with about 20% in the cytoplasmic fraction. The isolation of enzyme markers with the appropriate cellular compartment demonstrated that cytoplasmic, membranous and nuclear fractions can be isolated with minimal intercompartmental contamination using this fractionation scheme.

Nuclease Stability of Ribozymes and mRNA

The simplest and most sensitive way to monitor nuclease activity in cell fractions is to use end-labeled oligonucleotides. However, high levels of phosphatase activity in some biological extracts gives ambiguous results in nuclease experiments when '2P-5'-end-labeled oligonucleotides are used as substrates. To determine the phosphatase activity in the extracts, cellular fractions were incubated with cold ribozymes and trace amounts of 5'-end-labeled ribozyme in the presence of 1 mM Mg⁺² (or Zn⁺² with HeLa cytoplasmic extracts) to optimize digestion. After polyacrylamide gel electrophoresis of samples, digestion of the oligonucleotide was assessed both by staining and by autoradiography.

oligonucleotide basic Specifically, the digestion reaction contained substrate nucleic acid (an RNA oligonucleotide of 36 nucleotides) and cell fraction extract in a total volume of 100 µl. Aliquots (7 µl) were 25 taken after various periods of incubation at 37°C and added to 7 µl of gel loading buffer (95% formamide, 0.1% bromophenol blue, 0.1% xylene cyanol, and 20 mM EDTA). The samples were separated by electrophoresis on a $7\ \mathrm{M}$ Intact ribozymes were urea, 20% polyacrylamide gel. 30 visualized either by staining with Stains-all (United States Biochemical, Cleveland, OH), or autoradiography of 32P-labeled ribozyme. The stained gels and X-ray films were scanned on a Bio 5000 density scanner (U.S. Ribozymes were 5' end-labeled with T4 Biochemical). polynucleotide kinase (U.S. Biochemical) using 10 μCi of ^{32}P $\gamma\text{-ATP}$ (3,000 Ci/mmole, New England Nuclear, Boston, MA), and 20-25 pmoles of ribozymes. The unincorporated

nucleotides were separated from the product by G-50 spin chromatography. Nuclease assays contained 1-2 pmoles of oligonucleotides were ribozyme. All 32P-labeled Applied Biosystems 394 DNA/RNA synthesized on an synthesizer (Applied Biosystems Inc., Foster City, CA) according to manufacturer's protocols. The nuclear fractions were resuspended in a buffer containing 2.5 mM MgCl2. Experiments involving the nuclear fractions were performed in the presence of 1 mM Mg+2, or in combination 10 with 1 mM Mn+2, Ca+2, or Zn+2.

To measure the stability of mRNA, Vero cells were infected with herpes simplex virus (HSV) at a M.O.I. of 5 and total RNA was extracted (Chomczynski and Sacchi, 162 Anal. Biochem. 156, 1987). An RNase protection assay 15 was used to detect mRNA after incubation of total infected cellular RNA in cytoplasmic extracts. RNA probes were produced from PCR-amplified template DNA using T7 RNA polymerase (U.S. Biochemical) in the presence of ^{32}P $\alpha\text{-CTP}$ (3,000 Ci/mmole, New England Nuclear, Boston, MA). Template DNA was inactivated with 1 unit of RNase-free 20 DNaseI for 15 minutes at 37°C. Unincorporated nucleotides were removed by G-50 spin chromatography. Samples (6 μ 1) were taken from the nuclease assays after various periods of incubation at 37°C, added to 40 μl of 4 M GUSCN buffer (4 M guanidinium thiocyanate; 25 mM sodium citrate, pH 7; 0.5% sarcosyl; and 0.1 M 2-mercaptoethanol), and 5 μl of ^{32}P -labeled RNA probe (5x10 5 cpm/5 μ l, specific activity of Hybridization 1.8x106 cpm/µg) in 4 M GUSCN buffer. reactions were covered with mineral oil and incubated at 55°C for 12-16 hours, after which the hybridization reaction was mixed with 500 μ l of RNase buffer (0.4 M NaCl, 20 μ g/ml RNaseA, 2 units/ml Tl RNase) and incubated for 30 minutes at 37°C. RNase activity was quenched by incubation with 10 μl of 20% SDS and 10 μl of proteinase mg/ml), and the RNA was extracted using a phenol/chloroform mixture. The protected RNA fragment was purified by precipitation with an equal volume of

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isopropanol in the presence of 20 µg of carrier yeast tRNA. The RNA pellets were resuspended in gel loading buffer, heated to 95°C for 5 minutes and separated by electrophoresis on a 5% polyacrylamide, 7 M urea gel. Protected fragments were visualized by autoradiography, and the films were scanned with a Bio 5000 density scanner.

In experiments using these methods, the rate of digestion of ribozymes in Vero cell extracts was similar, demonstrating the lack of significant phosphatase activity in Vero cellular fractions. Similar results were observed with HeLa cellular fractions. In most extracts, ladders of digested fragments were observed; such ladders would not be expected if digestion was an artifact of phosphatase action. Thus, digestion using 5' end-labeled ribozymes is an accurate assessment of nuclease action in cellular extracts.

In other experiments, labeled ribozymes were incubated in various Vero and HeLa cellular fractions.

20 Incubation of ribozymes in either membranous or nuclear fractions resulted in a linear decrease of intact molecules over time. In contrast, no digestion of ribozymes occurred during a 24 hour incubation in Vero cytoplasmic extracts, and HeLa cytoplasmic extracts

25 exhibited a 20-30 minute delay in the onset of RNA digestion. After this refractory period, the rate of digestion was linear but not as rapid as the rates observed in any of the nuclear or membranous fractions.

The effect of four divalent cations (Mg⁺², Mn⁺², 30 Ca⁺², and Zn⁺²) on the nuclease activity of the cellular fractions was assessed. Vero cytoplasmic extracts were stimulated by the addition of 1 mM Mg⁺² or Mn⁺², while Ca⁺² or Zn⁺² had no effect. Nuclease activity in HeLa cytoplasmic extracts was enhanced only by the addition of 1 mM Zn⁺². Both Vero and HeLa membranous fractions exhibited maximum nuclease activity with the addition of Mg⁺² or Mn⁺² ions, while the addition of Ca⁺² significantly

reduced activity of the HeLa membranous fraction and abolished nuclease activity in the Vero membranous fraction. Addition of Zn⁺² to both membranous fractions resulted in a loss of all RNase activity. The Vero nuclear extract demonstrated roughly equivalent nuclease activity in the presence of either Mg⁺² alone or a Mg⁺² and Mn⁺² ion combination, less in the presence of Mg⁺² and Ca⁺², and no activity in the presence of Mg⁺² and Zn⁺². The effects of cation addition were not as dramatic with HeLa nuclear extracts. The nuclease activity of these fractions was greatest in the presence of Mg⁺² alone or Mg⁺² and Ca⁺² and decreased slightly with the addition of Mn⁺² or Zn⁺² to the Mg⁺² present in the extracts.

To verify that nuclease activity was dependent 15 upon added divalent cations, nuclease assays were performed using 1 mM Mg+2 in the presence and absence of 20 mM EDTA. For the HeLa cytoplasmic fractions, the Mg+2 was replaced with 1 mM Zn+2. The presence of 20 mM EDTA completely abolished nuclease activity in the Vero and 20 HeLa cytoplasmic fractions and Vero nuclear fractions. Nuclease activity in the HeLa membranous and nuclear fractions was partially inhibited by the addition of EDTA, while EDTA had no effect on the nuclease activity in the For comparative purposes, Vero membranous fraction. 25 reactions using DNA oligonucleotides were performed using different Vero fractions. All DNase activity in Vero cytoplasmic, membranous, and nuclear fractions was inhibited by 20 mM EDTA.

The stability of RNA oligonucleotides and HSV-1 mRNA were compared in the presence and absence of activity-enhancing divalent cations (1 mM Mg⁺², Vero cells; 1 mM Zn⁺², HeLa cells). Total cellular RNA from HSV-1 infected Vero cells (8 mg) and tracer amounts of ³²P-5'-end-labeled RNA oligonucleotides (1 pmole) were incubated with Vero or HeLa cytoplasmic extracts. In the absence of divalent cations, no substantial decrease of intact ribozymes was detected in assays, although mRNA was

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digested in both Vero and HeLa cytoplasmic extracts. After addition of divalent cations, digestion of ribozymes occurred in both Vero and HeLa cytoplasmic fractions. The rate of ribozyme digestion in HeLa extracts increased to levels similar to those observed with mRNA, while the rate of mRNA digestion remained greater than the rate of ribozyme digestion in Vero cytoplasmic fractions.

Thus, the stability of hammerhead ribozymes were compared in both Vero and HeLa cell cytoplasmic, membranous and nuclear fractions. Vero cytoplasmic and nuclear fractions were found to require Mg+2 for optimal nuclease activity, while the membranous fraction was not altered by the addition of divalent cations. membranous and nuclear fractions were also activated by Mg², while the cytoplasmic fractions required Zn² for nuclease activation. Relative stabilities of ribozymes and mRNAs were compared in Vero and HeLa cytoplasmic In the absence of appropriate divalent fractions. cations, little ribozyme digestion was observed in either 20 cytoplasmic preparation while mRNA was rapidly digested. The addition of Mg² to Vero cytoplasmic extracts and Zn² to the HeLa cytoplasmic extracts stimulated ribozyme degradation and enhanced mRNA digestion. These data show that the nuclease sensitivity of ribozymes is cell-type 25 specific, varies with the intracellular compartment studied and may not be able to be predicted from studies with mRNA. Notably, however, ribozymes appear stable in such cellular fractions for a period of time potentially sufficient to have a therapeutically useful activity.

30 Administration of Ribozyme

Selected ribozymes can be administered prophylactically, or to patients expressing mdr-1 mRNA, or having CML, leukemic conditions, Burkitt's lymphoma, follicular lymphoma, breast cancer, colon carcinoma, neuroblastoma, lung cancer, or pretumor cells, <u>e.g.</u>, by exogenous delivery of the ribozyme to an a desired tissue by means of an appropriate delivery vehicle, <u>e.g.</u>, a

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liposome, a controlled release vehicle, by use of iontophoresis, electroporation or ion paired molecules, or covalently attached adducts, and other pharmacologically approved methods of delivery. Routes of administration include intramuscular, aerosol, oral (tablet or pill form), topical, systemic, ocular, intraperitoneal and/or intrathecal. Alternatively, ribozymes may be administered to a tissue or afflicted cell ex vivo to eradicate tumorigenic cells prior to re-implantation (e.g., in the course of autologous bone marrow transplantation therapy). Expression vectors for immunization with ribozymes and/or delivery of ribozymes are also suitable.

. The specific delivery route of any selected ribozyme will depend on the use of the ribozyme. 15 Generally, a specific delivery program for each ribozyme will focus on unmodified ribozyme uptake with regard to intracellular localization, followed by demonstration of efficacy. Alternatively, delivery to these same cells in an organ or tissue of an animal can be pursued. Uptake studies will include uptake assays to evaluate cellular 20 ribozyme uptake, regardless of the delivery vehicle or Such assays will also determine strategy. intracellular localization of the ribozyme following uptake, ultimately establishing the requirements 25 maintenance of steady-state concentrations within the cellular compartment containing the target sequence (nucleus and/or cytoplasm). Efficacy and cytotoxicity can then be tested. Toxicity will not only include cell viability but also cell function.

- 30 Some methods of delivery that may be used include:
 - a. encapsulation in liposomes,
 - b. transduction by retroviral vectors,
 - c. conjugation with cholesterol,
 - d. localization to nuclear compartment utilizing nuclear targeting site found on most nuclear proteins,

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e. neutralization of charge of ribozyme by using nucleotide derivatives,

- f. use of blood stem cells to distribute ribozymes throughout the body, and
- g. electroporation.

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At least three types of delivery strategies are useful in the present invention, including: ribozyme modifications, particle carrier drug delivery vehicles, and retroviral expression vectors. Unmodified ribozymes, like most small molecules, are taken up by cells, albeit slowly. To enhance cellular uptake, the ribozyme may be modified essentially at random, in ways which reduce its charge but maintains specific functional groups. This results in a molecule which is able to diffuse across the cell membrane, thus removing the permeability barrier.

Modification of ribozymes to reduce charge is just one approach to enhance the cellular uptake of these larger molecules. The random approach, however, is not advisable since ribozymes are structurally and 20 functionally more complex than small drug molecules. The structural requirements necessary to maintain ribozyme catalytic activity are well understood by those in the art. These requirements are taken into consideration when designing modifications to enhance cellular delivery. The 25 modifications are also designed to reduce susceptibility to nuclease degradation. Both of these characteristics should greatly improve the efficacy of the ribozyme. Cellular uptake can be increased by several orders of magnitude without having to alter the phosphodiester linkages necessary for ribozyme cleavage activity.

Chemical modifications of the phosphate backbone will reduce the negative charge allowing free diffusion across the membrane. This principle has been successfully demonstrated for antisense DNA technology. The similarities in chemical composition between DNA and RNA make this a feasible approach. In the body, maintenance of an external concentration will be necessary to drive

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the diffusion of the modified ribozyme into the cells of the tissue. Administration routes which allow the diseased tissue to be exposed to a transient high concentration of the drug, which is slowly dissipated by systemic adsorption are preferred. Intravenous administration with a drug carrier designed to increase the circulation half-life of the ribozyme can be used. The size and composition of the drug carrier restricts rapid clearance from the blood stream. The carrier, made to accumulate at the site of infection, can protect the ribozyme from degradative processes.

Drug delivery vehicles are effective for both systemic and topical administration. They can be designed to serve as a slow release reservoir, or to deliver their contents directly to the target cell. An advantage of using direct delivery drug vehicles is that multiple molecules are delivered per uptake. Such vehicles have been shown to increase the circulation half-life of drugs which would otherwise be rapidly cleared from the blood stream. Some examples of such specialized drug delivery vehicles which fall into this category are liposomes, hydrogels, cyclodextrins, biodegradable nanocapsules, and bioadhesive microspheres.

From this category of delivery systems, 25 liposomes are preferred. Liposomes increase intracellular stability, increase uptake efficiency and improve biological activity.

Liposomes are hollow spherical vesicles composed of lipids arranged in a similar fashion as those lipids which make up the cell membrane. They have an internal aqueous space for entrapping water soluble compounds and range in size from 0.05 to several microns in diameter. Several studies have shown that liposomes can deliver RNA to cells and that the RNA remains biologically active.

For example, a liposome delivery vehicle originally designed as a research tool, Lipofectin, has

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been shown to deliver intact mRNA molecules to cells yielding production of the corresponding protein.

Liposomes offer several advantages: They are non-toxic and biodegradable in composition; they display long circulation half-lives; and recognition molecules can be readily attached to their surface for targeting to tissues. Finally, cost effective manufacture of liposome-based pharmaceuticals, either in a liquid suspension or lyophilized product, has demonstrated the viability of this technology as an acceptable drug delivery system.

Other controlled release drug delivery systems, such as nonoparticles and hydrogels may be potential delivery vehicles for a ribozyme. These carriers have been developed for chemotherapeutic agents and protein-based pharmaceuticals, and consequently, can be adapted for ribozyme delivery.

administration ribozymes of is Topical advantageous since it allows localized concentration at site of administration with minimal 20 adsorption. This simplifies the delivery strategy of the ribozyme to the disease site and reduces the extent of toxicological characterization. Furthermore, the amount of material to be applied is far less than that required Effective delivery for other administration routes. 25 requires the ribozyme to diffuse into the infected cells or through the skin to the underlying vasculature. Chemical modification of the ribozyme to neutralize is required for negative charge may be all that However, in the event that penetration. 30 neutralization is insufficient, the modified ribozyme can be co-formulated with permeability enhancers, such as Azone or oleic acid, in a liposome. The liposomes can either represent a slow release presentation vehicle in which the modified ribozyme and permeability enhancer 35 transfer from the liposome into the infected cell, or the liposome phospholipids can participate directly with the in enhancer modified and permeability ribozyme

facilitating cellular delivery. In some cases, both the ribozyme and permeability enhancer can be formulated into a suppository formulation for slow release.

Ribozymes may also be systemically administered. 5 Systemic absorption refers to the accumulation of drugs in the blood stream followed by distribution throughout the Administration routes which lead to systemic entire body. subcutaneous. intravenous, absorption include: intraperitoneal, intranasal, intrathecal and ophthalmic. Each of these administration routes expose the ribozyme to Subcutaneous tissue. accessible diseased administration drains into a localized lymph node which through the lymphatic network The rate of entry into the circulation has circulation. 15 been shown to be a function of molecular weight or size. The use of a liposome or other drug carrier localizes the ribozyme at the lymph node. The ribozyme can be modified to diffuse into the cell, or the liposome can directly participate in the delivery of either the unmodified or modified ribozyme to the cell. 20

A liposome formulation containing phosphatidylethanolomidomethylthiosuccinimide which can oligonucleotides to lymphocytes and macrophages is also useful for certain cancerous conditions. Furthermore, a 200 nm diameter liposome of this composition internalized as well as 100 nm diameter liposomes. 200 nm liposomes exhibit a 10-fold greater packaging capacity than the 100 nm liposomes and can accomodate larger molecules such as a ribozyme expression vector. 30 This oligonucleotide delivery system inhibits viral proliferation in these viruses that infect primary immune cells. This oligonucleotide delivery system prevents mRNA expression in affected primary immune cells. Whole blood studies show that the formulation is taken up by 90% of 35 the lymphocytes after 8 hours at 37°C. Preliminary biodistribution and pharmacokinetic studies yielded 70% of

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the injected dose/gm of tissue in the spleen after 1 hour following intravenous administration.

Intraperitoneal administration also leads to entry into the circulation with the molecular weight or size controlling the rate of entry.

Liposomes injected intravenously show accumulation in the liver, lung and spleen. The composition and size can be adjusted so that this accumulation represents 30% to 40% of the injected dose.

The remaining dose circulates in the blood stream for up to 24 hours.

The chosen method of delivery should result in cytoplasmic accumulation and molecules should have some nuclease-resistance for optimal dosing. Nuclear delivery 15 may be used but is less preferable. Most preferred delivery methods include liposomes (10-400 nm), hydrogels, microinjection polymers, controlled-release electroporation (for ex vivo treatments) and other pharmaceutically applicable vehicles. The dosage will 20 depend upon the disease indication and the route of administration but should be between 100-200 mg/kg of body weight/day. The duration of treatment will extend through the course of the disease symptoms, possibly continuously. The number of doses will depend upon disease delivery 25 vehicle and efficacy data from clinical trials.

within the cell is dependent upon the rate of uptake and degradation. Decreasing the degree of degradation will prolong the intracellular half-life of the ribozyme. Thus, chemically modified ribozymes, e.g., with modification of the phosphate backbone, or capping of the 5' and 3' ends of the ribozyme with nucleotide analogs may require different dosaging. Descriptions of useful systems are provided in the art cited above, all of which is hereby incorporated by reference herein.

The claimed ribozymes are also useful as diagnostic tools to specifically or non-specifically

detect the presence of a target RNA in a sample. That is, the target RNA, if present in the sample, will be specifically cleaved by the ribozyme, and thus can be readily and specifically detected as smaller RNA species.

The presence of such smaller RNA species is indicative of the presence of the target RNA in the sample.

Other embodiments are within the following claims.

Claims

- 1. An enzymatic RNA molecule which cleaves mRNA associated with development or maintenance of chronic myelogenous leukemia, promyelocytic leukemia, Burkitt's lymphoma or acute lymphocytic leukemia, follicular lymphoma, B-cell acute lymphocytic leukemia, breast cancer, colon carcinoma, neuroblastoma, and lung cancer, or which is active to specifically cleave mRNA expressed from a gene encoding multiple drug resistance.
- 2. The enzymatic RNA molecule of claim 1 which cleaves mRNA produced from the genes PML-RARA, C-myc, bcl-2, E2A-PRL, ErbB2/neu, ras, DCC, N-myc, L-myc or mdr-1.
- 3. The enzymatic RNA molecule of claim 1, which cleaves target mRNA having a sequence selected from 15 SEQ. ID. NOS. 1-9 in Fig. 2; SEQ. ID. NOS. 1-19 in Fig. 3; SEQ. ID. NOS. 1-62 in Fig. 4; SEQ. ID. NOS. 1-41 in Fig. 5; SEQ. ID. NOS. 1-22 in Fig. 6; SEQ. ID. NOS. 1-71 in Fig. 7; SEQ. ID. NOS. 1-118 in Fig. 8; SEQ. ID. NOS. 1-26 in Fig. 9; SEQ. ID. NOS. 1-66 in Fig. 10; and SEQ. ID. NOS. 1-17 in Fig. 11.
 - 4. The enzymatic RNA molecule of claims 1, 2 or 3, wherein said RNA molecule is in a hammerhead motif.
- 5. The enzymatic RNA molecule of claim 4, wherein said RNA molecule is in a hairpin, hepatitis Delta virus, group 1 intron, or RNaseP RNA motif.
 - 6. The enzymatic RNA molecule of claim 4, wherein said ribozyme comprises between 5 and 23 bases complementary to said mRNA.
- 7. The enzymatic RNA molecule of claim 6, 30 wherein said ribozyme comprises between 10 and 18 bases complementary to said mRNA.

- 8. A mammalian cell including an enzymatic RNA molecule of claims 1, 2 or 3.
- 9. The cell of claim 8, wherein said cell is a human cell.
- 10. An expression vector including nucleic acid encoding the enzymatic RNA molecule of claims 1, 2 or 3, in a manner which allows expression of that enzymatic RNA molecule within a mammalian cell.
- 11. A method for treatment of a disease caused

 10 by expression of an mdr-1 gene, chronic myelogenous
 leukemia, promyelocytic leukemia, Burkitt's lymphoma or
 acute lymphocytic leukemia, follicular lymphoma, B-cell
 acute lymphocytic leukemia, breast cancer, colon
 carcinoma, neuroblastoma, and lung cancer by administering

 15 to a patient an enzymatic RNA molecule of claims 1, 2 or
 3.
 - 12. The method of claim 11, wherein said patient is a human.

Cleavage site I I 5-AUUGGGGUCUGGAUA Therapeutic Substrate Ribozyme Segment CUG Α C-GAGU C-G G-C I G-C A G A A

FIG. 1

FIG. 2 (MDR-1 Gene)

NUCLEOTII NUMBER	DE SEQUENCE	•	
NT303	UCUUCCAAGCUCAAAGAAGCAGA	SEQ.ID.NO.	1
NT476	AAACUGAACAAUAAAAGUGAA	SEQ.ID.NO.	2
NT497	AAAGAUAAGAAGGAAAAG	SEQ.ID.NO.	3
NT514	GAAACCAACUGUCAGUGUA	SEQ.ID.NO.	4
NT546	GCUAUUCAAAUUGGCUUGACAA	SEQ.ID.NO.	5
NT585	GAACUUUGGC	SEQ.ID.NO.	6
NT612	CUGGACUUCC	SEQ.ID.NO.	7
NT641	GGAGAAAUGAC	SEQ.ID.NO.	8
NT686	CUGAUGUCAAACAUCACUAAUA	SEQ.ID.NO.	9



FIG. 3 (CML)

Nucleotide	Base	•
Reference <u>Number</u>	mRNA Target Sites	Seq. ID No.
bcr		
3135	CUGAUCUCCUCUGACUAUG	01
3196	GUGUUUCAGAAGCUU	02
3214	CCUGACAUCCGUGG	, 03
3226	GGAGCUGCAGAUGCUGACCA	04
3245	CAACUCGUGUGUGAAACUCC	05
3264	CAGACUGUCCACAGCAUUCCGCUGAC	.06
3282	CCGCUGACCAUCAAUAAGG	07
3333	CUGAAUGUCAUCGUCCA	08
3360	GAUUUAAGCAGAGUUCAA	0.9.
abl	·	
445	AGCCCUUCAGC	10
459	CAGUAGCAUCUG	11
488	CUGAGUGAA	12
542	GAAAAUGACCCC	13
554	AACCUUUUCGUUGC	14
571	GUAUGAUUUUGUG	15
592	AGAUAACACUCUAAGC	16
607	CAUAACUAAA	17
628	CCGGGUCUUAGGCUAUAAUCAC	18
647	CACAAUGGGGAAUGG	19

These targets are present in the 425 nucleotides surrounding the fusion sites of the bcr and abl sequences in the b2a2 and b3a2 recombinant mRNAs. Other sequences in the 5' portion of the bcr mRNA or the 3' portion of the abl mRNA may also be targeted for ribozyme cleavage. Cleavage at any of these sites in the fusion mRNA molecules will result in inhibition of translation of the fusion protein in treated cells.

FIG. 4 (PML)

Nucleotide Number Sequence	SEO.ID.NO.
1 CUCCCCUUCAGCUUCUCUCACGCACUCCAAGAUCUAA	ID.NO.01
78 ACCGCCGAUCUCCGAGGCCCCA	ID.NO.02
123 GGAGCCCACCAUGCCUCCCCCGAGACCC	ID.NO.03
179 GCCCCAGCCCUACAGAGCGAGCCCCCGCU	ID.NO.04
	ID.NO.05
251 CGGAAGCCAAGUGCCCGAAGCUGCUGCCU	ID.NO.06
289 ACGCUGUGCUCAGGA	ID.NO.07
346 CCCUGGCCCCUAG	ID.NO.08
380 AUAACGUCUUUUUCGAGAGUC	ID.NO.09
464 AAGAGUCGGCCGACUUCUGGUGCUUUGAGUGCGAG	ID.NO.10
522 CGAGGCACACCAGUGGUUCCUC 550 GAGGCCCGGCCCCUAGCAGAGCUGCGCAACCAGUC	ID.NO.11
591 UGAGUUCCUGGACGGCACCCGCAAGAC	ID.NO.12
618 CAACAACAUCUUCUGCUCCAACCCCAACCAC	ID.NO.13
655 CCUACGCUGACCAGCAUCUACUGCCGAGGAUGUUCCAA	
707 GUGCGCGCUCCUUGAC	ID.NO.15
737 AGCUCAAGUGCGACAUCAGCGCAGAGA	ID.NO.16
789 CGCCAUGACGCAGGCGCUGCAGGA	ID.NO.17
885 CGAGACCGAGGAGCUGAUCCGCGA	ID.NO.18
953 AGCUGCUGGAGGCUGUGGA	ID.NO.19
992 ACGAGGAGAUGGCC	ID.NO.20
1025 AUGCUGUGCUGCAGCGCAU	ID.NO.21
1069 AGGAUGAAĞUGCUA	ID.NO.22
1102 CUGGACAUGCACGGUUUC	ID.NO.23
1186 GAUGGCUUCGACGAGUUCAA	ID.NO.24
1235 UCACCCAGGGGAAAGCCAUUGAGACCCAGA	ID.NO.25
1277 AAGAGAUAGUGCCCA	ID.NO.26
1408 CGCCGCAGCAUCCA	ID.NO.27
1428 CAUGGUGUACACGU	ID.NO.28
1448 GGGACAAGAACUGCAUCAUCAACAA	ID.NO.29
1492 CAGUACUGCCGACUGCAGAAGUGCUUUGAAGUGGGC	ID.NO.30
1540 UCUGUGAGAAACGACCGAAACAAGAAGAA	ID.NO.31
1569 GAAGGAGGUGCCCAAGCCCGAGUGCUCU	ID.NO.32
1630 CUCAUUGAGAA	ID.NO.33
1690 AAAUACACUACGAACAACA	ID.NO.34
1735 AUUGACCUCUGGGACAA	ID.NO.35
1765 UCCACCAAGUGCAUCAUUAAGACUGUG	ID.NO.36
800 CAAGCAGCUGCCCGGCUUCACCACCCUCA	ID.NO.37
1829 CCAUCGCCGACCAGAUCACC	ID.NO.38



1854	CAAGGCUGCCUGCC	ID.NO.39
1894	ACGCGGUACACGC	ID.NO.40
1934	ACGGGCUGACCCUGAACCGGA	ID.NO.41
1955	CCCAGAUGCACAACGCUGGCUU	ID.NO.42
2018	UGCUGCCCUGG	ID.NO.43
2036	AUGAUGCGGAGACGGGGCUGCUCAGCG	ID.NO.44
2090	AGGACCUGGAGCAGCCGGACCG	ID.NO.45
2161	CGGAAGCGGAGGCCCAGCCCCCCA	ID.NO.46
2192	UCCCCAAGAUGCUAAUGAAGAUUACU	ID.NO.47
2356	GGUGGGGGGGGACGG	ID.NO.48
2391	GCCAGGCAGCUGUAGCCCCAGCCUCAGCC	ID.NO.49
2420	CCAGCUCCAACAGAAGCAGCCC	ID.NO.50
2450	ACUCCCGUGACCGCCACGCCACAUGGACA	ID.NO.51
2481	CAGCCCUCGCCCUCCG	ID.NO.52
2526	CAUGUGACCCCGCACCAG	ID.NO.53
2572	UACUGGGGACCUUCCC	ID.NO.54
2600	AGGGAGGAGCAGCGACUCCUUGGAC	ID.NO.55
2657	CCCACAGCCUGGGCUGACGUCAGA	ID.NO.56
2690	CAGGAACUGAG	ID.NO.57
2761	CCCUCUGCCCAGCUCACCACAUCUUCAUCACCAC	ID.NO.58
2824	CAGAACUCACAAGCCAUUGCUC	ID.NO.59
2864	ACCUCCCCCU	ID.NO.60
2953	AUUAAUUCUCGCUGGUUUUGUUUUUAUUUUAA	ID.NO.61
2995	тизатининима	ID.NO.62

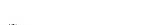
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FIG. 5 (c-Myc)

FIG. 5 (C-Myc)	
nucleotide mRNA target sequence	SEQ. ID. NO.
(469) CAGGACCCGCUUCUCUGAAAGGCUCUCCUU	ID.NO. 1
(553) GCGACGAUGCCCCUCAACGUUAGCU	ID.NO. 2
(589) AACUAUGACCUCGACUACGACUCGGU	ID.NO. 3
(629) ACUGCGACGAGGAGAACUUCUACCA	ID.NO. 4
(662) AGCAGCAGAGCGA	ID.NO. 5
(680) AGCCCCGGCGCCCAGCGAGGAUAUCUGGA	ID.NO. 6
(725) UGCCCACCCGCC	ID.NO. 7
(758) CCGGGCUCUGCUCGCCCUCCUA	ID.NO. 8
(783) UGCGGUCACACCCU	ID.NO. 9
(813) CAACGACGGCGGUGGCGGGGGGCUUCUCCACGGCCGACCA	ID.NO.10
(876) GGGAGGACAU	ID.NO.11
(907) UGCGACCCGGACGACGAGACCUUCAUCAAAAACAUCAUCA	ID.NO.12
(981) CGCCAAGCUCGUCUCAGAGAAGCUGGCCUCCUACCA	ID.NO.13
(1023) GCGCAAAGACAGCG	ID.NO.14
(1043) CGAACCCCGCCC	ID.NO.15
(1077) CUCCAGCUUGUACCUGCAGGA	ID.NO.16
(1114) UCAGAGUGCAUCGACCC	ID.NO.17
(1124) UCGACCCUCGGUGGUCUUCCCCUACCCUCUCAACGAC	ID.NO.18
(1168) UCGCCCAAGUCCUGCGCCUCGCAAGACUCCAGCGC	ID.NO.19
(1230) CUCCUCGACGGA	ID.NO.20
(1258) AGCCCCGAGCCCC	ID.NO.21
(1276) CUCCAUGAGGAGA	ID.NO.22
(1357) GUGGAAAAGAGG	ID.NO.23
(1376) CUGGCAAAAGGUCA	ID.NO.24
(1397) GAUCACCUUCUGCUGGAGGCCACAGCAAACCUCCUCACA	ID.NO.25
(1459) CACGUCUCCACACAUCAGCACAACUACGCA	ID.NO.26
(1496) CCUCCACUCGGAAGGACUAUCC	ID.NO.27
(1523) CCAAGAGGGUCAAGUUGGA	ID.NO.28
(1547) UCAGAGUCCUGAGACA	ID.NO.29
(1569) CAACAACCGAAAAUGCA	ID.NO.30
(1643) UGGAGCGCCAGAGG	ID.NO.31
(1662) CGAGCUAAAACGGAGCU	ID.NO.32
(1684) GCCCUGCGUGACCAGAUCCCGGA	ID.NO.33
(1712) AAAACAAUGAAAAGGCCCCCAAGGUAGUUAUCCUUAAAA	ID.NO.34
(1755) CACAGCAUACAUCCUGUCCGUCCAAG	ID.NO.35
(1786) GAGCAAAAGCUCAU	ID.NO.36
(1805) AAGAGGACUUGUUGCGGAAACGACGAGAACAGUUGAAAC	ID.NO.37
(1845) CAAACUUGAACAGCUACGGAACUCUUGUGCGU	ID.NO.38
(1883) AAGUAAGGAAAACGA	ID.NO.39

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(1904)	CUAACAGAAAUGUCCUGAGCAAUCACCUAUGAACU	ID.NO.40
(1964)	ACCUCACAACCU	ID.NO.41

FIG. 6 (BCL-2)

NUCLEOT	TIDE mRNA TARGET SEQUENCE	SEQ.ID.	NO.
(1359)	AAAUAGCUGGAUUAUAACU	ID.NO.	01
(1415)	GGCGAGAGGUGCCGUUGCCUUUUCCUCU	ID.NO.	02
(1504)	AUGAAGUACAUC	ID.NO.	03
(1522)	AAGCUGUCGCAGAGGGGCUACGA	ID.NO.	04
(1592)	CACCGGGCAUCUUCUCCUCCCAGCCCGGGCACA	ID.NO.	05
(1656)	CGCCAGGACCUCGCCGCUGCAG	ID.NO.	06
(1719)	CCCGGUGCCACCUGUG	ID.NO.	07
(1729)	CCUGUGGUCCACCUGGCCCUCCGCCAAG	ID.NO.	80
(1683)	GGCUGCCCCGGC	ID.NO.	09
(1742)	UGGCCCUCCGCCAAGCC	ID.NO.	10
(1807)	AGCCAGCUGCACCUGACGC	ID.NO.	11.
(1836)	GCGGGGACGCUUUGCCACG	ID.NO.	12
(1937)	AGAGCGUCAAC	ID.NO.	13
(1972)	AACAUCGCCCUGUGGAUGA	ID.NO.	14
(1997)	ACCUGAACCGGCACCUGCACACCUGGAUCCAGGAUA	ID.NO.	15
(2043)	GGAUGCCUUUGUGGAACUGUACGGCCCCAGCAUG	ID.NO.	16
(2081)	CUCUGUUUGAUUUCUCCUGGCUGUCUC	ID.NO.	17
(2100)	CUGUCUCUGAAGACUCU	ID.NO.	18
(2135)	UGGGAGCUUGCAUCACC	ID.NO.	19
(2180)	GUCAACAUGCCUGCCCCAAACAAAUAUGCAAAAGG	ID.NO.	20
(2235)	AAUAUGCAUUGUCAGUGAUGUACCAUGAAACAAA	ID.NO.	21
(2262)	AAACAAAGCUGCAGGCU	ID.NO.	22

Number	Sequence	SEO.ID.NO.
25	CAGCAGGGUUUCCAGGCCUGA	ID.NO.01
68	GAAUGAACCAGC	ID.NO.02
110	AGGAGCUCAGUGACCUCCUGGACUUCAGCAUGAUG	ID.NO.03
151	CUGCCUGUCACCAA	ID.NO.04
187	CUGGCCGGGCGCAGUUC	ID.NO.05
211	UCAGGUCUUGAGGA	ID.NO.06
297	CGAGGGCACCCACUUCA	ID.NO.07
319	UCGCACAGCAGCCUCUCUUCAUCCACAUUCCUGGGA	ID.NO.08
425	GCCUGACUCAGGCUGGCUUCCUGUCAGG	ID.NO.09
453	CGAGCUGGCCCUCAACAGCCCCGGGCCCCU	ID.NO.10
507	CUCCCAGUACUACCCCUCCUACUCCGGCA	ID.NO.11
536	GCUCCCGGCGGAGAGCGGCAG	ID.NO.12
557	ACGGCAGCCUAGACACGCAGCCCAAGAAGGUCCGGAAG	ID.NO.13
610	CCAUCCUCGGUGUACCCACCCAGCUCAGGUG	ID.NO.14
701	CCGCCCCUUCUACGUGGCA	ID.NO.15
726	CAGCCUGCACCC	ID.NO.16
814	CUCCCGCCGGUAGCGGCCCGGU	ID.NO.1
844	AGUGGAAGCAGCAC	ID.NO.18
866	GUGGCCUGCACCAGCACGAGCGUAUGGGCUA	ID.NO.19
904	CAUGGAGCAGAGGUGAACGGUGG	ID.NO.20
927	GCUCCCAUCUGCAUCCUCCUUCUCCUCAGCCC	ID.NO.2
957	CCGGAGCCACGUAC	ID.NO.23
.021	CUGGGCUCCGAGGGACCA	ID.NO.2
047	CAGCUCCGGGAUGCCCUCGGCAAAGCACUGGCCUC	ID.NO.2
108	AAUAACUUCUCGUCC	ID.NO.2
L153	GGCCUGGCAGGAACGUCACA	ID.NO.2
L196	GUGCCUUAUCGCCCAGCUACGA	ID.NO.2
L224	UCUCCACGGCCUGCAGAGUAAGAUAGAAGACCACCUGGA	ID.NO.2
L272	CCACGUGCUCC	ID.NO.2
L325	UGCCUGGCCACGGGGC	ID.NO.3
1346	CCUCAGGUUUCA	ID.NO.3
1377	CGGGCGGCACGCAGGC	ID.NO.3
1416	GGACGGCCUCGCAGGCAGCACCAGCCUCAUGCACAAC	ID.NO.3
1453	CACGCGGCCCUCC	ID.NO.3
1474	CCAGGCACCCUCCCUGACCUGUCUCG	ID.NO.3
1506	CGACUCCUACAGUGUUUUGAGUAUCCGA	ID.NO.3
1551	ACCCACAGACCCCCAGCUGAUGC	ID.NO.3
1606	GCGGGCCUGAGAAGGCCGGA	ID.NO.3



1632	GGCGGCAGCGGCAGCGGCGGCUUCUG	ID.NO.39
1677	AGACAACUCAGUGGAGCAUUCAGAUUACA	ID.NO.40
1719	ACAGAUCAGACAAAUCUACCAUACG	ID.NO.41
1744	GAGCUGGAGAAAUACGAGCAGGCCUGCAA	ID.NO.42
1773	CGAGUUCACCACCCACGUGAUGAAUCUCCUGCGAG	ID.NO.43
1845	UGAGCGGAUGGUCAGCAUCAUCCACCGCAA	ID.NO.44
1892	UGCAGCUCAAGCAGAGCACGUGC	ID.NO.45
1926	GAUCCUGCGUUCCCGAUUUCUGGAUGCGCGGCG	ID.NO.46
1972	UUCAACAAGCAAGCGAC	ID.NO.47
1995	CCUGAAUGAAU	ID.NO.48
2017	CAUCUCAGCAAC	ID.NO.49
2045	AAGCCAAAGAGGAGUUAGCCAAGAAGUGUGGC	ID.NO.50
2088	CCAGGUAUCAAACUGG	ID.NO.51
2135	ACAUAGGUAAAUUUCAAGAGGAAG	ID.NO.52
2159	CCAAUAUUUAUGCUGCCAA	ID.NO.53
2178	AACAGCUGUCACUGCUACCAAUGU	ID.NO.54
2209	CAUGGAAGCCAAGCUAACUCGCCCUCAACUCCCAACUC	ID.NO.55
2259	CAGUUCUUUUAACAUGUCAAACU	ID.NO.56
2295	CAUGAGCGUGCAGUCACUCAAUGGGGAUUCUUACCAA	ID.NO.57
2347	GCCAACGUGCA	ID.NO.58
2381	AUGUUAUCAGCCAGACAGGAGGAUACA	ID.NO.59
2418	CGCAGCCAGUCAGAUGUAC	ID.NO.60
2442	GCAGGGCAUCAGUGCUAAU	ID.NO.61
2468	GGCAGGAUGCUACUACCCCUUCAUCAGU	ID.NO.62
2503	CCUACAGAAGGCCCUGGCA	ID.NO.63
2546	GAUCUCCCAGCAAUCGCAUCCCGG	ID.NO.64
2638	ACUGGAGGUCGAAGCAAUC	ID.NO.65
2695	GGGAUGCUAUUUCAGCCAAUCU	ID.NO.66
2722	CUUCUUUAUACUCUCUUCCCUUUUUUUUUUU	ID.NO.67
2758	AAGCCACCCUUCC	ID.NO.68
2777	CCAGCUGUCAGCCUGGUUUUCGUCAUCUUCC	ID.NO.69
2808	CUGCCCUGUGCCUC	ID.NO.70
2900	CAAAAAAUUACAAAGAAAAUAAUAAAA	ID.NO.71

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FIG. 8 (Breast Cancer)

	-10. 0 (110000 001.001)	
Nucleot	tide mRNA target sequence	SEQ.ID.NO.
(67)	CCUUUACUGCGCCGCGC	ID.NO. 01
(94)	CACCCUCGCAGCACCCCGC	ID.NO. 02
(113)	GCCCCGCGCCCCCAGCCGGGUCCAGCCGGAGCCA	ID.NO. 03
(159)	AGCCGCAGUGAGCACCAUGGAGCU	ID.NO. 04
(240)	CACCCAAGUGUGCACCGGCACAGACAUGAAGCU	ID.NO. 05
(285)	CAGUCCCGAGAC	ID.NO. 06
(305)	ACAUGCUCCGCCACCUCUACCAGGGCUG	ID.NO. 07
(369)	GCCCACCAAUGCCAGCCUGUC	ID.NO. 08
(413)	UGCAGGGCUACGU	ID.NO. 09
(449)	GGCAGGUCCCACUG	ID.NO. 10
(468)	GCUGCGGAUUGUGCGAGGCACCCAGCU	ID.NO. 11
(500)	AGGACAACUAUGCCCUGGCCGUGCUAGACAAUGGAGAC	ID.NO. 12
(580)	GGCCUGCGGAGCUGCAGCCUCACAGAGA	ID.NO. 13
(714)	UCUCACACUGAUAGACACCAACCGCUCUCGGGCCUGC	ID.NO. 14
(747)	CUGCCACCCUGUUCUCCGAUGUGUAAGGG	ID.NO. 15
(794)	AGAGUUCUGAGGAUUGUCAGA	ID.NO. 16
(849)	CCGCUGCAAGGGGCCACUGCCCACUG	ID.NO. 17
(893)	GUGCUGCCGGCU	ID.NO. 18
(916)	AAGCACUCUGACUGCCUGCCUCCACUUCAAC	ID.NO. 19
(951)	CACAGUGGCAUC	ID.NO. 20
(969)	GCUGCACUGCCCAGCC	ID.NO. 21
(990)	CACCUACAACACAGACACGUUUGAG	ID.NO. 22
(1025)	AUCCCGAGGGCCGGUA	ID.NO. 23
(1075)	UACAACUACCUUUCUACGGACGU	ID.NO. 24
(1103)	CCUGCACCCUCGUCUGCCCCUGCACAACC	ID.NO. 25
(1132)	AAGAGGUGACAGCAGGAUGGA	ID.NO. 26
(1218)	GCACUUGCGAG	ID.NO. 27
(1235)	GGGCAGUUACCAGUGCCAAUAU	ID.NO. 28
(1262)	AGUUUGCUGGCUGCAAGAAGAUCUUUG	ID.NO. 29
(1294)	CUGGCAUUUCUGCCGGAGAGCUUUGAUGGGGAC	ID.NO. 30
(1327)	CCAGCCUCCAACACUGCCCC	ID.NO. 31
. (1331)	CCUCCAACACUGCCCC	ID.NO. 32
(1357)	GAGCAGCUCCAAGUGUUUGAGAC	ID.NO. 33
(1392)	CACAGGUUACCUAUAC	ID.NO. 34
	AGCAUGGCCGGACAGCCUGACCUCAGCGUC	ID.NO. 35
· ·	ACCUGCAAGUAAUC	ID.NO. 36
•	CACAAUGGCGCCUACUC	ID.NO. 37
(1505)	CCCUGCAAGGGCUGGGCA	ID.NO. 38

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		•		
	(1538)	UGCGCUCACUGAGGGAA	ID.NO.	39
	(1560)	CAGUGGACUGGCCCUCAUCCACCAUAACAC	ID.NO.	40
	(1590)	CCACCUCUGCUUCGUGCACACGGUGCCCUGGGA	ID.NO.	41
	(1622)	CCAGCUCUUUCGGAACC	ID.NO.	42
	(1662)	UGCCAACCGGCCAGAGGACGA	ID.NO.	43
	(1699)	CUGGCCUGCCACCAGCU	ID.NO.	44
	(1749)	CACCCAGUGUCAACUGC	ID.NO.	45
	(1782)	GGGCCAGGAGUGCG	ID.NO.	46
	(1804)	UGCCGAGUACUGCA	ID.NO.	47
	(1832)	AGUAUGUGAAUGCCA	ID.NO.	48
	(1892)	CAGUGACCUGUUUUGGACC	ID.NO.	49
	(1916)	CUGACCAGUGUGGCCUGUGCCCACUAUAAGGAC	ID.NO.	50
	(1986)	GAAACCUGACCUCCUACAUGCCAU	ID.NO.	51
•	(2033)	AGGGCGCAUGCCAG	ID.NO.	52
	(2055)	CAUCAACUGCACCCACUCCUGUGUGGA	ID.NO.	53
	(2123)	CUCUGACGUCCAUCAUCUCUGCGGU	ID.NO.	54
	(2163)	GGUCGUGGUCUUG	.ID.NO.	55
	(2212)	AAGAUCCGGAAGUACACGAUGCGGAGA	ID.NO.	56
	(2249)	AGGAAACGGAGC	ID.NO.	57
	(2286)	AGCGAUGCCCAA	ID.NO.	58
	(2303)	CGCAGAUGCGGAUCCUGAAAGAGACGGAGCUGAGGAAGG	ID.NO.	59
	(2375)	UCUACAAGGGCAUCUGGA	ID.NO.	60
	(2405)	AGAAUGUGAAAAUUCCAG	ID.NO.	61
	(2432)	AAGUGUUGAGGGAAAACACAUCCCCCAAAGCCAA	ID.NO.	62
	(2466)	CAAAGAAAUCUUAGACGAAGCAUACG	ID.NO.	
	(2516)	AUGUCUCCGCCUUCUGGGCA	ID.NO.	64
	(2542)	CUGACAUCCACGGUGCAGCUGGUGACAC	ID.NO.	65
	(2576)	UGCCCUAUGGCUGCCUCUUAGACCAU	ID.NO.	66
	(2630)	CCCAGGACCUGCUGAACUGGUGUAU	ID.NO.	
	(2660)	UUGCCAAGGGGA	ID.NO.	
	(2678)	ACCUGGAGGAUGUGC	ID.NO.	
	(2716)	GCUCGGAACGUGCU	ID.NO.	
	(2735)	AGAGUCCCAACCAUGU	ID.NO.	
	(2675)	GCUACCUGGAGGAUGU	ID.NO.	
	•	GCUCGGAACGUGCU	ID.NO.	
	•	AGUCCCAACCAUGUCAAAAUUACAGACUUC	ID.NO.	
		CUCGGCUGCUGGACAUUGACGAGACAGAGUACCAUGCAG	ID.NO.	
		AAGUGGAUGGCG	ID.NO.	•
		GUCCAUUCUCCGCCG	ID.NO.	
	(2870)	UCACCCACCAGAGUGAUGUGGGAGUUAUGGUGUG	ID.NO.	/ 6

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GUGGGAGCUGAUGACU	ID.NO. 79
GGCCAAACCUUAC	ID.NO. 80
ACCUGCUGGAAAAGGGGGAGCG	ID.NO. 81
AGCGGCUGCCCAGC	ID.NO. 82
AUCUGCACCAUUGAUGUCUACAUGAUCAU	ID.NO. 83
GAAUGUCGGCCAAGAUUCCG	ID.NO. 84
CCCCAGCGCUUUGUGGUCAUCCAGAAUGAGGACUUGGGC	ID.NO. 85
UGGACACCUUCU	ID.NO. 86
UCACUGCUGGAGGAC	ID.NO. 87
CAUGGGGACCUGGUGGA	ID.NO. 88
AGGAGUAUCUGGUACCC	ID.NO. 89
GGGCUUCUUCUGUCCAGACCC	ID.NO. 90
UGGUCCACCACAGGCACCGCAGC	ID.NO. 91
UACCAGGAGUG	ID.NO. 92
ACACUAGGGCUGGAGCCCUCUGAAGAGGAGGCCCC	ID.NO. 93
CUGGCUCCGAU	ID.NO. 94
AUGGUGACCUGGGAA	ID.NO. 95
AGCCAAGGGGCUGCAAAGCCUCCCCACACAUGACCCC	ID.NO. 96
AGCCCUCUACAGC	ID.NO. 97
AGUGAGGACCCCACA	ID.NO. 98
CCUGCCCUCUGAGACUGAUGGCUACGUUGCCCCCCUGA	ID.NO. 99
GCCUGAAUAUGUGAACCAGC	ID.NO. 100
UCGGCCCCAGC	ID.NO. 101
CCCCGAGAGGGCCCUCUGCCUGCUGCCCGACCUGCUGG	ID.NO. 102
CUGGAAAGGCCCAAGACUCUCU	ID.NO. 103
UCGUCAAAGACGUUUUUG	ID.NO. 104
AGUACUUGACACCCCAGGGAGGAGCUGCCCCUCAGCCC	ID.NO. 105
CCUUCGACAACCUCUAUUA	ID.NO. 106
GACACCUACGGC	ID.NO. 107
ACGUGCCAGUGUGAA	ID.NO. 108
ACCCAAGUCC	ID.NO. 109
AGCAGGGAAGGCCUGA	ID.NO. 110
GAGGGCCCUCCGA	ID.NO. 111
CAGGGGAACCUGCCAUGCCAGGAACCUGUCCUA	ID.NO. 112
CCCAAUGAGACUC	ID.NO. 113
UCCUUCCAGAU	ID.NO. 114
AAGCCUUAGGGAAGCUGGCCUGAGAGGGGAAGCGGCCC	ID.NO. 115
GUGUCUAAGAACAAAAGCGACCCAUUCAGAGACUG	ID.NO. 116
GAAACCUAGUACUGCCCCCAUGAGGAAGGAACAGCAA	ID.NO. 117
UUAGUUUUUACUUUUUUUUUUUUUUUUUUUUUUUAAAG	ID.NO. 118
	GUGGGACUGAUGACU GGCCAAACCUUAC ACCUGCUGGAAAAGGGGGAGCG AGCGGCUGCCCCAGC AUCUGCACCAUUGAUGUCUACAUGAUCAU GAAUGUCGGCCAAGAUUCCG CCCCAGCGCUUUGUGGUCAUCAGAAUGAGGACUUGGGC UGGACAGCACUUCU UCACUGCUGGAGAC CAUGGGGGACCCCC GGGCUUCUUCUGUCCAGAAUGAGGACUUGGCC GGGCUUCUUCUGUCCAGACCC UGGUCCACCACAGGCACCCC UGGUCCACCACAGGCACCCC UGGCUCCGAU AUGGGGGGGCCCUCUGAAGAGGAGGCCCC CUGGCUCCGAU AUGGUGACCCCACACAUGACCC AGCCACAGGCACCCCC AGCCCUCUACAGC AGUGAGGACCCCACACAUGACCCC CUGGCCCCCACACACAUGACCCC AGCCCUCUACAGC CCUGCACUCUACAGC CCUGCACUCUAGAGACCGCCCCCCCCCC

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FIG. 9 (Colon Carcinoma)

Nucleotide	Target mRNA sequence	ID. N	os.
(1) UCCUAGGCGGCG	·	ID.NO.	01
(18) GGCGGCGGAGGCAG	CAGCG	ID.NO.	02
(42) GGCAGUGGCGGC	GG	ID.NO.	03
(68) CGGCUCGGCCAG	UACUCCCGGCCC	ID.NO.	04
(120) GCGCAGGCACUG	AAGGCGGCGGCGGGCCAGAGGCUCAG	ID.NO.	05
(203) AUAAACUUGUGG	UAGUUGGAGCUUGUGGCGU	ID.NO.	06
(248) UGACGAUACAGC	UA	ID.NO.	07
(275) UUGUGGACGAAU	AUGAUCCAACAAUA	ID.NO.	80
(307) UCCUACAGGAAG	CAAGUAGUAAUUGAU	ID.NO.	09
(374) AAGAGGAGUACA	GUGCAAUGAGGGA	ID.NO.	10
(435) AUUUGCCAUAAA	UAAUACUAAAUCAUUUGAA	ID.NO.	11
(466) GAUAUUCACCAU	UAUAGAGAACAAAUUAAA	ID.NO.	12
(511) GAAGAUGUACCU	AUGGUCCUAGUAGGAAAUAAAUGU	ID.NO.	13
(547) GAUUUGCCUUCU	AGAACAGUAG	ID.NO.	14
(602) AUGGAAUUCCU		ID.NO.	15
(631) AAGACAAGACAG	egg .	ID.NO.	16
(675) AGAAAUUCGAA		ID.NO.	17
(686) AACAUAAAGAAA	AGAUGAGCAAAGAUGGUAAAAAGA	ID.NO.	18
(722) AGAAAAAGAAGU	ICAAAGACAAAGUGUGUA	ID.NO.	19
(781) UAAGGCAUACUA	AGU	ID.NO.	20
(820) ACUAAAUUAUU	AG	ID.NO.	21
(840) UUUAGCAUUAC	AAU	ID.NO.	. 22
(863) CUGCUCCAUGCI	AGACUGUUAGC	ID.NO	
(907) AAAAUGACAGU	GGA ·	ID.NO	
(946) UAUUCCCAGAGI	סט	ID.NO	
(970) ACUAGCAAUGC	CUGU	ID.NO	. 26

FIG. 10 (Neuroblastoma)

Nucleotid Number	mRNA Sequence	SEO.ID.NO.
1	ACAGUCAUCUGUCUGGACGC	ID. NO. 01
39	GCUCCUGGGAACUGUGUUGGAGCCGAGCAA	ID. NO. 02
84	AAGCGCGCACAGACUGUAGCCAUCCGAGGACA	ID. NO. 03
153	GAAUCGCCUCCGGAUCC	ID. NO. 04
257	GACGGGAUUGCGACG	ID. NO. 05
339	UUUCCCAGAAAAGCCAGUUCCAG	ID. NO. 06
367	AAGGCAUCCUGGCU	ID. NO. 07
387	GACCCGCCCUAA	ID. NO. 08
434	UCUGCGAAAAGAAAUUCCCU	ID. NO. 09
460	UAGAAGAUCUGUCUGUGUUU	ID. NO. 10
535	CCACGGGAAGGAAGCACCCCCGGUAUUAAAACG	ID. NO. 11
577	GAAAGAAGCCCUCAGUCGCCGGC	ID. NO. 12
635	ACCAUGCCGGGCAUGAUCUGCAAGAAC	ID. NO. 13
662	CCAGACCUCGAGUUUGACUCGCUACAGC	ID. NO. 14
690	CCUGCUUCUACCC	ID. NO. 15
708	AAGAUGACUUCUACUUCGGCGGCC	ID. NO. 16
750	GGGAGGACAUCUGGAAGAAGUUUGA	ID. NO. 17
785	ACGCCCCGCUGUC	ID. NO. 18
847	CACGGAGAUGCUGCUUGAGAACGAGCUGUG	ID. NO. 19
923	GGCCUCACCCCCA	ID. NO. 20
953	GACUGCAUGUGGA	ID. NO. 21
976	CGCCCGCGAGAAGCUGGAGCGCCCCGUGAGC	ID. NO. 22
1049	ACCGCCAGUCCCCG	ID. NO. 23
1074	CCGCCAGCCCUGCGGGUC	ID. NO. 24
1133	GCCCUGCCCGCCGAGCUCGCCCACCC	ID. NO. 25
1159	GGCCGCCGAGUGCGUGGAUC	ID. NO. 26
1191	UCCCCUUUCCCGUGAACAAGCGCGAGCCAGCG	
1234	AGCCCCGGCCAGUGCCCCGGCG	ID. NO. 28
	CAGUACCUCCGGA	ID. NO. 29
	ACCCUGAGCGAUUCAGAAUG	ID. NO. 30
	ACAUGAGAGUAACUAG	ID. NO. 31
1441	CAGAUGAUGAAGAUGA	ID. NO. 32
1471	AAGAGGAAGAAAUCGAC	ID. NO. 33
	GUUCCUCCUCAACACCAA	ID. NO. 34
	UCACCACAUUCACCAUCACUGUGCGUCCCAAGA	ID. NO. 35
	UCCUCAAACGAU	
1629	AUCCACCAGCAGCACAACUAUGCCGCCCC	ID. NO. 3/

1744	CCCCAAAGGCUAAGAGCU	ID.	NO.	39
1767	CCCCGAAACUCUGACU	ID.	NO.	40
1789	ACAGUGAGCGUCGCAGAAACCACAACAUCCUGG	ID.	NO.	41
1847	GUCCAGCUUUCUCAC	ID.	NO.	42
1889	AAAGAAUGAGAAGGCC	ID.	NO.	43
1917	AUUUUGAAAAA	ID.	NO.	44
1935	GAGUAUGUCCACUCCCUCCAGGCC	ID.	NO.	45
2031	CACGCUCGGACUUGCUAGACGCUUCUCAAAA	ID.	NO.	46
2055	UCAAAACUGGACAGUCACUGCCACUUUGCACAU	ID.	NO.	47
2115	GUGUUGACAUUAAG	ID.	NO.	48
2162	GAGUUCGGCUCUG	ID.	NO.	49
2222	AGCCUGCAUCCCAGGA	ID.	NO.	50
2271	UCCAUGACAGCGCUAAACGUU	ID.	NO.	51
2297	CGGUUGGGAGCCUCU	ID.	NO.	52
2321	GAAGUCACCUUGUGUGUUCCAAGUUUCCAAACA	ID.	NO.	53
2376	UAAAAUGGUGCUUAAGUUCCAGCAGAUGCCACA	ID.	NO.	54
2417	UUGCCAUUUGAU	ID.	NO.	55
2435	GGGGAACAUUU	ID.	NO.	56
2453	UACCAUUGACACAU	ID.	NO.	57
2478	AUACAUCCUGGGUA	ID.	NO.	58
2527	AAGUUCAUACCUAAGUACUGU	ID.	NO.	59
2670	UUUGAUACUUAUAUU	ID.	NO.	60
2713	UGAGUAGAUAUUACUUUAUCACUUUUUGAACUA	ID.	NO.	61
2760	AGAAAUUUACUAUAUAUAUAUGCCUUUUUCCUA	ID.	NO.	62
2818	UGUUCAUGUUU	ID.	NO.	63
2877	AAUGUAUAUUU		NO.	
2916	AUACCUCAUGUUUAUGAAAAU		NO.	
2942	GCUUAAUUAAA	ID.	NO.	66

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FIG. 11 (L-myc)

nucleotide		
number	mRNA target sequence	Seq.ID. Nos.
9	UGCAGCUCGCGCUCCCGCGCCGAUCCCGAGA	ID.NO. 01
61	AGCGAGGGGGGG	ID.NO. 02
93	GCUCGUGAGUGC	ID.NO. 03
152	GCUGCCCGAG	ID.NO. 04
181	GCUCCAGGUGGC	ID.NO. 05
200	UGGAGCGAGGGAGCGACAUGGACUACGACUC	ID.NO. 06
249	ACGACUAUGACUGCGGG	ID.NO. 07
266	GAGGAUUUCUACCGCUCCACGGC	ID.NO. 08
289	GCCCAGCGAGGACAUCUGGA	ID.NO. 09
327	CAUCGCCCCCACGUC	ID.NO. 10
391	UGGUCCCCGGAGCCGUGGCCCG	ID.NO. 11
414	GAGGGUGCACCGGAGA	ID.NO. 12
445	GGGCCACUCGAAAGGCU	ID.NO. 13
465	AGGAACUACGCCUCCAUCAUACGCC	ID.NO. 14
490	GUGACUGCAUGUGGAGCGGCUUCUCGGCCCGGG	ID.NO. 15
525	AACGGCUGGAGA	ID.NO. 16
557	GCUCCUGGCGCGC ·	ID.NO. 17

INTERNATIONAL SEARCH REPORT

international application No. PCT/US93/04573

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A. CL	ASSISTED ATTOM OF STREET		
IPC(5)	ASSIFICATION OF SUBJECT MATTER :A61K 31/70; C07H 19/10; C12N 5/16, 15/63		
US CL	:514/44; 536/23.1; 435/240.2, 320 1		
According	to International Patent Classification (IPC) or to be	oth national classification and IPC	
B. FIE	LDS SEARCHED		
Minimum	documentation searched (classification system follo-	wed by classification symbole)	
U.S. :	514/44; 536/23.1; 435/240.2, 320.1	ay moday	•
Documenta	ation searched other than minimum documentation to	the extent that such documents are include	d in the fields searched
			e in the fields scattlied
Electronic	data base consulted during the international search	(name of data base and, where practicable	s. search terms used)
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C. DO	CUMENTS CONSIDERED TO BE RELEVANT		
			/
Category*	Where		Relevant to claim No.
\mathbf{Y}	WO, A1, 91/18624 (Scalon) 12	December 1991, see entire	1-12
	document.	, 444	
Y	WO 11 01 1000	•	
1	WO, A1, 91/18625 (Scalon) 12	December 1991, see entire	1-12
	document.		
Y	WO A1 02/00000 m ::		
1	WO, A1, 92/00080 (Reddy, et. al.)	09 January 1992, see entire	1-12
	document.		
Y	Call Volume 47 : 110 0		
	Cell, Volume 47, issued 10 October	1986, M. L. Cleary, et. al.,	I-12
	"Cloning And Structural Analysis (or cDNAs For bcl-2 And A	
	Hybrid bcl-2/Immunoglobulin Trant (14;18) Translocation", pages 19-28,	script Resulting From The	
1	, pages 19-20,	see entire document.	
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X Furthe	er documents are listed in the continuation of Box (
	cial categories of cited documents: ument defining the general state of the art which is not considered	"T" later document published after the inter date and not in conflict with the applica	national filing date or priority
	e part or particular relevance .	principle or theory underlying the inve	ntion
	ier document published on or after the international filing date	"X" document of purticular relevance; the considered novel or cannot be considered to the considered t	claimed invention cannot be
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7.00	ALLEICABLE	Telephone No. (703) 308-0196	•

INTERNATIONAL SEARCH REPORT

International application No. PCT/US93/04573

Category*		
	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim
Y	Science, Volume 247, issued 09 March 1990, N. Sarver, et. al., "Ribozymes As Potential Anti-HIV-1 Therapeutic Agents", pages 1222-1225, see entire document.	1-12
?	Mol. Cell. Biol., Volume 8, No. 1, issued January 1988, F. Kaye, et. al., "Structure And Expression Of The Human L-myc Gene Reveal A Complex Pattern Of Alternative mRNA Processing", pages 186-195, see entire document.	I-12
7	Nature, Volume 304, issued 11 August 1983, J. P. McGrath, et. al., "Structure And Organization Of The Human Ki-ras Proto-oncogene And A Related Processed Pseudogene", pages 501-506, see entire document.	1-12
	Proc. Natl. Acad. Sci. USA, Volume 82, issued October 1985, K. Semba, et. al., "A v-erbB-Related Protoocogene, C-erB-2, Is Distinct From The c-erbB-1/Epidermal Growth Factor-Receptor Gene And Is Amplified In A Human Salivary Gland Adenocarcinoma", pages 6497-6501, see entire document.	I-12
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BOX II. OBSERVATIONS WHERE UNITY OF INVENTION WAS LACKING This ISA found multiple inventions as follows:

- I. Claims 1-12, drawn to an enzymatic RNA which cleaves mRNA associated with chronic myelogenous leukemia, a cell containing this RNA, a vector that encodes this RNA, and a method of use in treating a disease.
- II. Claims 1-12, drawn to an enzymatic RNA which cleaves mRNA associated with promyelocytic leukemia, a cell containing this RNA, a vector that encodes this RNA, and a method of use in treating a disease.
- III. Claims 1-12, drawn to an enzymatic RNA which cleaves mRNA associated with Burkitt's lymphoma, a cell containing this RNA, a vector that encodes this RNA, and a method of use in treating a disease.
- IV. Claims 1-12, drawn to an enzymatic RNA which cleaves mRNA associated with acute lyphocytic leukemia, a cell containing this RNA, a vector that encodes this RNA, and a method of use in treating a disease.
- V. Claims 1-12, drawn to an enzymatic RNA which cleaves mRNA associated with follicular lymphoma, a cell containing this RNA, a vector that encodes this RNA, and a method of use in treating a disease.
- VI. Claims 1-12, drawn to an enzymatic RNA which cleaves mRNA associated with B-cell acute lymphocytic leukemia, a cell containing this RNA, a vector that encodes this RNA, and a method of use in treating a disease.
- VII. Claims 1-12, drawn to an enzymatic RNA which cleaves mRNA associated with breast cancer, a cell containing this RNA, a vector that encodes this RNA, and a method of use in treating a disease.
- VIII. Claims 1-12, drawn to an enzymatic RNA which cleaves mRNA associated with colon carcinoma, a cell containing this RNA, a vector that encodes this RNA, and a method of use in treating a disease.
- IX. Claims 1-12, drawn to an enzymatic RNA which cleaves mRNA associated with neuroblastoms, a cell containing this RNA, a vector that encodes this RNA, and a method of use in treating a disease.
- X. Claims 1-12, drawn to an enzymatic RNA which cleaves mRNA associated with lung cancer, a cell containing this RNA, a vector that encodes this RNA, and a method of use in treating a disease.